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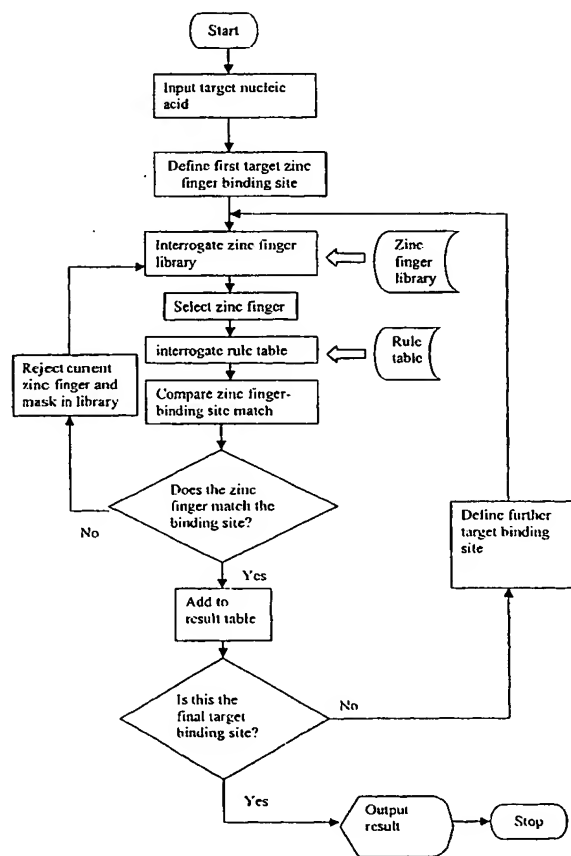
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(54) Title: COMPOSITE BINDING POLYPEPTIDES



(57) Abstract: Disclosed herein are polypeptides with novel DNA binding specificities, constructed from combinations of zinc fingers, and methods for their preparation and use.

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COMPOSITE BINDING POLYPEPTIDES**TECHNICAL FIELD**

5 The present disclosure is in the fields of molecular biology and protein design; in particular, the design of sequence-specific binding proteins for regulation of gene expression.

10 BACKGROUND

Protein-nucleic acid recognition is a commonplace phenomenon that is central to a large number of biomolecular control mechanisms that regulate the functioning of eukaryotic and prokaryotic cells. For instance, protein-DNA interactions form the basis of the 15 regulation of gene expression and are thus one of the subjects most widely studied by molecular biologists.

A wealth of biochemical and structural information explains the details of protein-DNA recognition in numerous instances, to the extent that general principles of recognition 20 have emerged. Many DNA-binding proteins contain independently folded domains for the recognition of DNA, and these domains in turn belong to a large number of structural families, such as the leucine zipper, the "helix-turn-helix" and zinc finger families.

Despite the great variety of structural domains, the specificity of the interactions observed 25 to date between protein and DNA most often derives from the complementarity of the surfaces of a protein α -helix and the major groove of DNA. See, e.g., Klug, (1993) Gene 135:83-92. In light of the recurring physical interaction of α -helix and major groove, the tantalising possibility arises that the contacts between particular amino acids and DNA bases could be described by a simple set of rules; in effect a stereochemical recognition 30 code which relates protein primary structure to binding-site sequence preference.

It is clear, however, that no code will be found which can describe DNA recognition by all DNA-binding proteins. The structures of numerous complexes show significant differences in the way that the recognition α -helices of DNA-binding proteins from different structural families interact with the major groove of DNA, thus precluding 5 similarities in patterns of recognition. The majority of known DNA-binding motifs are not particularly versatile, and any codes which might emerge would likely describe binding to a very few related DNA sequences.

Even within each family of DNA-binding proteins, moreover, it has hitherto appeared 10 that the deciphering of a code would be elusive. Due to the complexity of the protein-DNA interaction, there does not appear to be a simple "alphabetic" equivalence between the primary structures of protein and nucleic acid which specifies a direct amino acid to base relationship.

15 International patent application WO 96/06166 addresses this issue and provides a "syllabic" code that explains protein-DNA interactions for zinc finger nucleic acid binding proteins. A syllabic code is a code that relies on more than one feature of the binding protein to specify binding to a particular base, the features being combinable in the forms of "syllables", or complex instructions, to define each specific contact. Segal, 20 D. J., Dreier, B., Beerli, R. R. & Barbas, C. F. (1999) Proc. Natl. Acad. Sci. USA 96, 2758-2763 present a method of constructing zinc fingers polypeptides, based on 16 individual zinc finger domains which bind sequences of the form 5'-GXX-3', where X is any base. See also U.S. Patent No. 6,140,081. The latter method has the severe 25 limitation that it does not provide instructions permitting the specific targeting of triplets containing nucleotides other than G in the 5' position of each triplet, which greatly restricts the potential target sequences of such generated zinc finger peptides.

International patent application WO98/53057 addresses the above problems by 30 recognizing that zinc fingers can specify overlapping 4 bp subsites, and therefore synergy between adjacent zinc finger domains is an important consideration in selecting zinc finger nucleic acid-binding domains to specifically target any sequence.

With the recent completion of the human genome project and the rapidly advancing fields of transgenic animals and plants, thousands of uncharacterised (and characterised) genes have (and will) become valid targets for functional genomics and other such projects.

Concomitantly, 'designer' zinc finger peptides are emerging as one of the most universal 5 and desirable ways of regulating the expression of specific genes within cells. See, for example, Choo, Y., Sanchez-Garcia, I. & Klug, A. (1994) *Nature* 372: 642-645; Beerli, R. R., Dreier, B. & Barbas, C. F. III (2000) *Proc. Natl. Acad. Sci. USA* 97: 1495-1500; Kim, J-S. & Pabo, C. O. (1998) *Proc. Natl. Acad. Sci. USA* 95: 2812-2817; Kang, J. S. & Kim, J-S. (2000) *J. Biol. Chem.* 275: 8742-8748; Zhang *et al.* (2000) *J. Biol. Chem.* 10 275:33,850-33,860; Liu *et al.* (2001) *J. Biol. Chem.* 276:11,323-11,334; and Ren *et al.* (2002) *Genes. Devel.* 16:27-32. See also WO 00/41566 and WO 01/19981. Hence, a rapid method of creating multi-zinc finger peptides for the up- or down-regulation of any specific gene is highly desirable.

As stated above, synergy between adjacent zinc finger peptides is an important factor in 15 specific DNA recognition. Moreover, the findings reported in co-owned WO 01/53480, which is hereby incorporated by reference, demonstrate that poly-zinc finger peptides constructed from strings of 2-finger domains can provide greater DNA binding specificity.

20 Traditional strategies of zinc finger mutagenesis and selection, such as phage display, particularly if employed for the selection of 2-zinc finger units to target any desired binding site are limited by the size of the library that can be cloned into host/vector systems, such as phage. Due to limitations in library size imposed by such constraints, it is impossible to include an exhaustive combination of randomisations to cover all 25 potentially important sequence-space. Furthermore, for important applications of engineered zinc finger peptides, such as for gene therapy or transgenic animal systems, engineered zinc finger peptides run the significant risk of eliciting a harmful immunological reaction in the host animal.

30 The human genome sequencing project has also revealed the presence of almost 700 endogenous zinc finger-containing proteins. Assuming that each of these proteins

contains at least 2 finger modules, there are probably at least 2,000 natural zinc finger modules in the human genome alone. Similar numbers are expected in other animal and plant genomes.

5 SUMMARY

The present invention recognises the potential importance of designer zinc finger peptides in therapeutic and transgenic applications in animals and plants. Furthermore the present invention acknowledges that the safety of such applications is of primary importance.

10

The present invention provides the isolation of natural zinc finger modules, from genomes such as human, mouse, chicken, arabidopsis and other species, and the construction of non-natural combinations of such zinc finger modules, to create multi-finger domains, and to provide and determine novel nucleic acid binding specificities.

15

Such a procedure will allow the identification of the novel zinc finger domains that bind any desired nucleic acid sequence, particularly sequences of between 6 and 10 nucleotides long. The first advantage of such technology is that millions of years of natural evolution, to create specific nucleotide-binding zinc finger modules, are captured to create novel nucleic acid-binding domains. Also, use of poly-zinc finger peptides constructed from such units for targeted gene regulation avoids the potentially harmful effects of host immune responses. The present invention thus greatly enhances the possibilities for the use of zinc finger transcription factors for *in vivo* applications, such as gene therapy and transgenic animals.

20

25

In a first aspect, therefore, there is provided a composite binding polypeptide comprising a first natural binding domain derived from first natural binding polypeptide, and a second natural binding domain derived from a second natural binding polypeptide, wherein said first and second natural binding polypeptides may be the same or different; which polypeptide binds to a target, said target differing from the natural target of the both the first and the second binding polypeptides.

Preferably, said first and second natural binding polypeptides are different polypeptides.

Binding polypeptides according to the invention comprise two or more natural binding domains, advantageously three or more natural binding domains; advantageously, six or more domains are included. These are preferably arranged in a 3x2 conformation, 5 separated by linker sequences.

The binding domains are preferably nucleic acid binding domains, and the composite polypeptide is preferably a nucleic acid binding polypeptide. Most preferably, the composite polypeptide is a zinc finger polypeptide, and the natural binding domains are 10 zinc finger domains.

Zinc finger binding domains can comprise any type of zinc finger or zinc-coordinated structure including, but not limited to, Cys2-His2 (SEQ ID NO:1) zinc finger binding domain or Cys3-His (SEQ ID NO:2) zinc finger binding domains.

15 In a further aspect, there is provided a library of natural binding domains. The natural binding domains are the domains that may be assembled into polypeptides according to the previous aspect of the invention. Preferably, the library is of natural zinc finger nucleic acid binding domains.

20 Said zinc finger domains may comprise a linker attached thereto. Any linker amino acid sequence known in the art can be used. Advantageously, the linker comprises the amino acid sequence TGEKP (SEQ ID NO:3).

25 In a further aspect, the invention provides a method for selecting a binding polypeptide capable of binding to a target site, comprising:

- (a) providing a library of natural binding domains;
- (b) assembling two or more of said domains to form a composite polypeptide;
- (c) screening said composite polypeptide against the target site in order to

30 determine its ability to bind the target site.

Preferably, the natural binding domains are zinc finger binding domains.

Furthermore, the invention provides methods for designing a composite binding polypeptide, comprising:

- (a) providing information defining a target site;
- 5 (b) selecting, from a database of natural binding domains, a sequence of binding domains, separated by linker sequences, which is predicted to bind to the target site;
- (c) displaying the sequence of binding domains and linkers and optionally assembling the binding polypeptide from a library of said domains.

10 In certain embodiments, the binding domains are zinc finger domains. In certain embodiments, a binding domain sequence that will bind a particular target site is predicted by the application of one or more rules that define target binding interactions for the binding domains. In additional embodiments, a nucleotide sequence encoding the binding domains is assembled and introduced into a cell such that the composite binding 15 polypeptide is expressed.

In one embodiment, zinc fingers can be considered to bind to a nucleic acid triplet, in which case domains can be selected according to one or more of the following rules:

- (a) if the 5' base in the triplet is G, then position +6 in the α -helix is Arg; or 20 position +6 is Ser or Thr and position ++2 is Asp;
- (b) if the 5' base in the triplet is A, then position +6 in the α -helix is Gln and ++2 is not Asp;
- (c) if the 5' base in the triplet is T, then position +6 in the α -helix is Ser or Thr and position ++2 is Asp;
- 25 (d) if the 5' base in the triplet is C, then position +6 in the α -helix may be any amino acid, provided that position ++2 in the α -helix is not Asp;
- (e) if the central base in the triplet is G, then position +3 in the α -helix is His;
- (f) if the central base in the triplet is A, then position +3 in the α -helix is Asn;
- (g) if the central base in the triplet is T, then position +3 in the α -helix is Ala, Ser 30 or Val; provided that if it is Ala, then one of the residues at -1 or +6 is a small residue;

(h) if the central base in the triplet is C, then position +3 in the α -helix is Ser, Asp, Glu, Leu, Thr or Val;

(i) if the 3' base in the triplet is G, then position -1 in the α -helix is Arg;

(j) if the 3' base in the triplet is A, then position -1 in the α -helix is Gln;

5 (k) if the 3' base in the triplet is T, then position -1 in the α -helix is Asn or Gln;

(l) if the 3' base in the triplet is C, then position -1 in the α -helix is Asp.

In a further embodiment, the zinc fingers can be considered to bind to a nucleic acid quadruplet and domains can be selected according to one or more of the following rules:

10 (a) if base 4 in the quadruplet is G, then position +6 in the α -helix is Arg or Lys;

(b) if base 4 in the quadruplet is A, then position +6 in the α -helix is Glu, Asn or Val;

(c) if base 4 in the quadruplet is T, then position +6 in the α -helix is Ser, Thr, Val or Lys;

15 (d) if base 4 in the quadruplet is C, then position +6 in the α -helix is Ser, Thr, Val, Ala, Glu or Asn;

(e) if base 3 in the quadruplet is G, then position +3 in the α -helix is His;

(f) if base 3 in the quadruplet is A, then position +3 in the α -helix is Asn;

(g) if base 3 in the quadruplet is T, then position +3 in the α -helix is Ala, Ser or

20 Val; provided that if it is Ala, then one of the residues at -1 or +6 is a small residue;

(h) if base 3 in the quadruplet is C, then position +3 in the α -helix is Ser, Asp, Glu, Leu, Thr or Val;

(i) if base 2 in the quadruplet is G, then position -1 in the α -helix is Arg;

(j) if base 2 in the quadruplet is A, then position -1 in the α -helix is Gln;

25 (k) if base 2 in the quadruplet is T, then position -1 in the α -helix is His or Thr;

(l) if base 2 in the quadruplet is C, then position -1 in the α -helix is Asp or His;

(m) if base 1 in the quadruplet is G, then position +2 is Glu;

(n) if base 1 in the quadruplet is A, then position +2 Arg or Gln;

(o) if base 1 in the quadruplet is C, then position +2 is Asn, Gln, Arg, His or Lys;

30 (p) if base 1 in the quadruplet is T, then position +2 is Ser or Thr.

In a preferred embodiment, zinc fingers are considered to bind to a nucleic acid quadruplet and domains are selected according to one or more of the following rules:

- (a) if base 4 in the quadruplet is G, then position +6 in the α -helix is Arg; or position +6 is Ser or Thr and position ++2 is Asp;
- 5 (b) if base 4 in the quadruplet is A, then position +6 in the α -helix is Gln and ++2 is not Asp;
- (c) if base 4 in the quadruplet is T, then position +6 in the α -helix is Ser or Thr and position ++2 is Asp;
- 10 (d) if base 4 in the quadruplet is C, then position +6 in the α -helix may be any amino acid, provided that position ++2 in the α -helix is not Asp;
- (e) if base 3 in the quadruplet is G, then position +3 in the α -helix is His;
- (f) if base 3 in the quadruplet is A, then position +3 in the α -helix is Asn;
- 15 (g) if base 3 in the quadruplet is T, then position +3 in the α -helix is Ala, Ser or Val; provided that if it is Ala, then one of the residues at -1 or +6 is a small residue;
- (h) if base 3 in the quadruplet is C, then position +3 in the α -helix is Ser, Asp, Glu, Leu, Thr or Val;
- (i) if base 2 in the quadruplet is G, then position -1 in the α -helix is Arg;
- (j) if base 2 in the quadruplet is A, then position -1 in the α -helix is Gln;
- 20 (k) if base 2 in the quadruplet is T, then position -1 in the α -helix is Asn or Gln;
- (l) if base 2 in the quadruplet is C, then position -1 in the α -helix is Asp;
- (m) if base 1 in the quadruplet is G, then position +2 is Asp;
- (n) if base 1 in the quadruplet is A, then position +2 is not Asp;
- 25 (o) if base 1 in the quadruplet is C, then position +2 is not Asp;
- (p) if base 1 in the quadruplet is T, then position +2 is Ser or Thr.

25

Two or more composite polypeptides comprising two or more domains which are selected for binding to two or more target sites can be combined to provide a composite polypeptide which binds to an aggregate binding site comprising the two or more target binding sites.

30

In a still further aspect, the invention provides a computer-implemented method for designing a zinc finger polypeptide that binds to a target nucleic acid sequence, comprising the steps of:

- (a) providing a system comprising at least storage means for storing data relating to a library of zinc fingers; storage means for storing a rule table; means for inputting target nucleic acid sequence data; processing means for generating a result; and means for outputting the result;
- 5 (b) inputting sequence data for a target nucleic acid molecule;
- (c) defining a first target zinc finger binding site in said nucleic acid molecule;
- 10 (d) interrogating the zinc finger library and rule table storage means, comparing zinc fingers to the target zinc finger binding site according to the rule table and selecting zinc finger data identifying a zinc finger capable of binding to said target site;
- (e) defining at least one further target zinc finger binding site and repeating step (d); and
- 15 (f) outputting the selected zinc finger data.

Such a method may further comprise sending instructions to an automated chemical synthesis system to assemble a zinc finger polypeptide as defined by the zinc finger data obtained in (f).

20 In additional embodiments, the sequence of one or more oligonucleotides encoding a composite binding polypeptide can be determined from the sequence of a composite binding polypeptide, and the one or more oligonucleotides can be synthesized by any number of well-known methods.

25 Preferably, a composite binding polypeptide is tested for binding to a target sequence, and data from said testing is used to select, from a plurality of possibilities, a composite binding polypeptide that binds with optimal affinity and specificity to the target site.

30 Advantageously, two or more zinc finger polypeptides are combined to form a zinc finger polypeptide capable of binding to an aggregate binding site comprising two or more target sites.

The rule table preferably comprises rules as set forth above.

BRIEF DESCRIPTION OF THE FIGURES

5 **Figure 1** shows a flowchart depicting part of the logic used in the selection of zinc fingers from a natural library in accordance with the invention. The logic set forth in Figure 1 may be supplemented, for example using Rules relating to zinc finger overlap. Functional testing of zinc fingers for binding to the desired binding site may be implemented in an automated fashion and integrated with the zinc finger design system.

10

Figure 2 is a schematic representation of the human zinc finger mini-library construction procedure. Synthetic zinc finger coding oligonucleotides are assembled into full-length ds expression constructs by overlap PCR.

15 **Figure 3** is a schematic representation of the fluorescent ELISA assay used to detect zinc finger peptides bound to double stranded DNA target sites. Streptavidin (7), biotinylated DNA target (5) linked to biotin (6), 3-finger peptide (4) fused to HA-tag (3), anti-HA antibody (2) fused to horseradish peroxidase (HRP, 1).

20 **Figure 4** depicts ELISA scores of 384 library 2 constructs screened against the 5'-GCG-TGG-GCG-3' (SEQ ID NO:4) target site. Six constructs showed significant binding, and are termed C8, G16, I19, I23, J19 and K19, according to their coordinates on the 384-well plate.

25 **Figure 5** depicts ELISA scores of selected library 2 members; B10, C8, G16, I23, J19, and K19, against different DNA target sites. The sequences of the target sites are (from back of graph to front): 5'-GCG-TGG-GCG-3' (SEQ ID NO:5) ; 5'-CCA-CTC-GGC-3' (SEQ ID NO:6); 5'-CCT-AGG-GGG-3'(SEQ ID NO:7); 5'-GGA-TAA-GCG-3' (SEQ ID NO:8); 5'-GGG-AGG-CCT-3' (SEQ ID NO:9); 5'-GCG-TAA-GGA-3' (SEQ ID NO:10); 5'-GCG-GGG-GGA-3' (SEQ ID NO:11); and no DNA control (front row).

Figure 6 depicts a schematic representation of the 3-zinc finger library constructed according to the procedure described in Example 2.

DETAILED DESCRIPTION

5

Definitions

Unless defined otherwise, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art (e.g., in cell culture,

10 molecular genetics, nucleic acid chemistry, hybridisation techniques and biochemistry).

The practice of the present invention will employ, unless otherwise indicated, conventional techniques of chemistry, molecular biology, microbiology, recombinant DNA, immunology, chemical methods, pharmaceutical formulations and delivery and treatment of patients, which are within the capabilities of a person of ordinary skill in the

15 art. Such techniques are explained in the literature. See, for example, J. Sambrook, E. F. Fritsch, and T. Maniatis, 1989, *Molecular Cloning: A Laboratory Manual*, Second

Edition, Books 1-3, Cold Spring Harbor Laboratory Press; Ausubel, F. M. et al. (1995 and periodic supplements; *Current Protocols in Molecular Biology*, ch. 9, 13, and 16, John Wiley & Sons, New York, N.Y.); B. Roe, J. Crabtree, and A. Kahn, 1996, *DNA*

20 *Isolation and Sequencing: Essential Techniques*, John Wiley & Sons; J. M. Polak and James O'D. McGee, 1990, *In Situ Hybridisation: Principles and Practice*; Oxford

University Press; M. J. Gait (Editor), 1984, *Oligonucleotide Synthesis: A Practical Approach*, IRL Press; and, D. M. J. Lilley and J. E. Dahlberg, 1992, *Methods of*

Enzymology: DNA Structure Part A: Synthesis and Physical Analysis of DNA Methods in Enzymology, Academic Press. Each of these general texts is herein incorporated by reference.

30 The term "library" is used according to its common usage in the art, to denote a collection of different polypeptides or, preferably, a collection of nucleic acids encoding different polypeptides. The libraries of natural zinc finger peptides referred to herein comprise or encode a repertoire of polypeptides of different sequences, each of which has a preferred binding sequence.

The terms "polypeptide", "peptide" and "protein" are used interchangeably to refer to a polymer of amino acid residues, preferably including naturally occurring amino acid residues. Artificial amino acid residues are also within the scope of the invention, but the 5 exclusive use of naturally-occurring amino acids is preferred in order to maintain the natural nature of the binding domains. There are 20 common amino acids, each specified by a different arrangement of three adjacent DNA nucleotides by the genetic code. These are the building blocks of proteins. Joined together in a strictly ordered chain by peptide bonds, the sequence of amino acids determines each polypeptide molecule. The 20 10 common amino acids are: alanine, arginine, aspartic acid, glutamic acid, glutamine, glycine, histidine, isoleucine, leucine, phenylalanine, proline, serine, threonine, tryptophan, tyrosine, valine, cysteine, methionine, lysine, and asparagine. Virtually all of these amino acids (except glycine) possess an asymmetric carbon atom, and thus are potentially chiral in nature.

15

As used herein, "nucleic acid" includes both RNA and DNA, and nucleic acids constructed from natural nucleic acid bases or synthetic bases, or mixtures thereof. Modified nucleic acids such as, for example, PNAs and morpholino nucleic acids, are also included in this definition.

20

A "gene", as used herein, is the segment of nucleic acid (typically DNA) that is involved in producing a polypeptide chain or ribonucleic acid gene product. It includes regions preceding and following the coding region (leader and trailer) as well as intervening sequences (introns) between individual coding segments (exons). Preferably, "gene" 25 includes the necessary control sequences for gene expression, as well as the coding region encoding the gene product.

A "binding polypeptide" is a polypeptide capable of binding to a specific target. Although, as is well known, polypeptides are capable of non-specific binding to a wide 30 range of substrates, it is also known that certain polypeptides, such as antibodies and other members of the immunoglobulin superfamily, zinc fingers, leucine zipper polypeptides, peptide aptamers and the like can bind specifically to target sites or

molecules. Generally, specific binding is preferably achieved with a dissociation constant (K_d) of 100 μ M or lower; preferably 10 μ M or better; preferably 1 μ M or better; and ideally 0.5 μ M or better. Binding polypeptides can be nucleic acid binding polypeptides which bind to nucleic acid in a target sequence-specific manner, such as zinc finger polypeptides. Unless specifically noted, no difference is intended herein between terms such as "peptide", "polypeptide" and "protein".

A "natural binding polypeptide" is a binding polypeptide encoded by the genome of a living organism such as, for example, a plant or animal.

10

A "composite" polypeptide is a polypeptide that is assembled from a plurality of components. In a preferred embodiment, the invention provides composite binding polypeptides that are assembled from a plurality of individual natural binding domains as set forth in detail herein. Typically, such domains are zinc finger nucleic acid binding domains.

15

A "natural binding domain" (or module) is a domain of a naturally occurring polypeptide that is capable of specific binding to a target as defined above. The terms "domain" and "module", according to their ordinary signification in the art, refer to a discrete continuous part of the amino acid sequence of a polypeptide that can be equated with a particular function. Protein domains or modules are largely structurally independent and can retain their structure and function in different environments. In certain embodiments, a natural binding domain or module is a zinc finger that binds a triplet or quadruplet nucleotide sequence.

20

Preferably, each of the individual natural binding domains that make up a composite binding polypeptide contain no changes in sequence, as compared to the natural sequence. However, those skilled in the art will understand that certain changes including conservative amino acid substitutions, as well as additions or deletions, may be made without altering the function of a domain. Moreover, where the changes are consistent 30 with sequences common to the species from which the domain is derived, such as for

example being present in consensus sequences, they are unlikely to give rise to immunological problems.

Conservative amino acid substitutions may be made, for example according to Table 1.

5 Amino acids in the same block in the second column and preferably in the same line in the third column may be substituted for one another:

Table 1

ALIPHATIC	Non-polar	G A P
		I L V
	Polar - uncharged	C S T M
		N Q
AROMATIC	Polar - charged	D E
		K R
AROMATIC		H F W Y

A domain is "derived" from a protein if it is effectively removed from a naturally-occurring protein for use in a composite binding polypeptide. Removal may be physical

5 removal, by cleavage of the protein; more commonly, however, the sequence of the domain is determined and the domain is synthesised by protein synthesis techniques to be a copy of the naturally-occurring domain. Alternatively, a nucleic acid encoding the domain is synthesized and expressed in a cell. *In vitro* synthesised domains, or *in vitro* synthesized polynucleotides encoding naturally-occurring domains, are considered to be 10 "derived" from the natural protein if they recapitulate the sequence of the naturally-occurring domain.

A "target" is a molecule or part thereof to which a binding polypeptide or a binding doamin is capable of specific binding. The "natural target" of a binding polypeptide is 15 the target to which that polypeptide binds in nature; *e.g.*, in a living cell. In the case of zinc finger polypeptides, for instance, the natural target is the nucleotide sequence to which the polypeptide binds in a living cell. Sequences other than the natural target, as defined herein, to which a zinc finger polypeptide may bind *in vitro* are not natural targets.

20

In the case of nucleic acid binding polypeptides, therefore, the term "target" may be substituted or supplemented with "binding site" or "binding sequence." Where binding sites are assembled to form larger binding sites, which are bound by multi-domain

binding polypeptides, such binding sites are referred to as "aggregate binding sites", indicating that they are formed by the juxtaposition of two or more individual binding sites. The aggregate binding sites can comprise contiguous individual binding sites, or individual binding sites interspersed by one or more intervening nucleotides or sequence
5 of nucleotides.

The present invention relates to naturally-occurring zinc fingers and their use as specific nucleic acid binding modules in combinations not present in nature. This invention provides methods of determining and/or predicting the nucleotide binding specificities of
10 natural zinc finger modules. Also provided are methods of constructing poly-zinc finger peptides containing at least one natural zinc finger module, from libraries of natural zinc finger peptides, and methods of screening such peptides to determine their preferred nucleotide binding specificity. Moreover, the invention provides for the use of combinations of such natural zinc finger modules in poly-zinc finger peptides not present
15 in nature, to bind any desired nucleotide sequence.

Poly-zinc finger peptides of this invention may contain 2, 3, 4, 5, 6 or more zinc finger modules. Natural zinc finger modules of this invention may preferably be linked by canonical, flexible or structured linkers, as set out below and in WO 01/53480, the disclosure of which is hereby incorporated by reference. More preferably, the linkers are
20 canonical linkers such as -TGEKP- (SEQ ID NO:3).

The poly-zinc finger peptides of this invention can be given useful biological functions by the addition of effector domains, creating chimeric zinc finger peptides. Preferably, such chimeric zinc finger peptides may be used to up- or down-regulate desired genes, *in vitro* or *in vivo*. Preferable effector domains include transcriptional repressor domains, transcriptional activator domains, transcriptional insulator domains, chromatin remodelling domains, enzymatic domains, and signalling / targeting sequences or domains. To cause a desired biological effect composite binding polypeptides can bind to one or more suitable nucleotide sequences *in vivo* or *in vitro*. Preferred DNA regions
25 from which to effect the up- or down-regulation of specific genes include promoters, enhancers or locus control regions (LCRs). Other suitable regions within genomes,
30

which may provide useful targets for composite binding polypeptides include telomeres and centromeres.

The expression of many genes is also achieved by controlling the fate of the associated

5 RNA transcript. RNA molecules often contain sites for RNA-binding proteins, which determine RNA half-life. Hence, composite binding polypeptides can also control endogenous gene expression by specifically targeting RNA transcripts to either increase or decrease their half-life within a cell.

10 Composite binding polypeptides can also be fused to epitope tags, which can be detected by antibodies, and may therefore be used to signal the presence or location of a particular nucleotide sequence in a mixed pool of nucleic acids, or immobilised on the surface of a chip or other such surface.

15 Intracellular localization of composite binding polypeptides can be regulated, for example, by fusion to a localization domain, for example, a nuclear localization sequence or a localization domain as disclosed, for example, in PCT/US01/42377.

20 **a. Nucleic Acid Binding Polypeptides**

This invention preferably relates to nucleic acid binding polypeptides. Preferably, the binding polypeptides of the invention are DNA binding polypeptides. Particularly preferred examples of nucleic acid binding polypeptides are zinc finger peptides.

25 Zinc finger peptides typically contain strings of small nucleic acid binding domains, each stabilised by the co-ordination of zinc. These individual domains are also referred to as "fingers" and "modules". A zinc finger recognises and binds to a nucleic acid triplet, or an overlapping quadruplet, in a DNA target sequence. However, zinc fingers are also known to bind RNA and proteins. Clemens, K. R. *et al.*, (1993) *Science* 260: 530-533; Bogenhagen, D.F. (1993) *Mol. Cell. Biol.* 13: 5149-5158; Searles, M. A. *et al.*, *J. Mol. Biol.* 301: 47-60 (2000); Mackay, J. P. & Crossley, M. (1998) *Trends Biochem. Sci.* 23: 1-4.

Preferably, there are 2 or more zinc fingers, for example 2, 3, 4, 5, 6, or 7 zinc fingers, in each zinc finger polypeptide. Advantageously, there are 3 or more zinc fingers in each zinc finger polypeptide.

5

All of the DNA binding residue positions of zinc finger peptides, as referred to herein, are numbered from the first residue in the α -helix of the finger, ranging from +1 to +9. “-1” refers to the residue in the framework structure immediately preceding the α -helix in a zinc finger peptide. Residues referred to as “++” are residues present in an adjacent

10 (C-terminal) peptide. Where there is no C-terminal adjacent peptide, “++” interactions do not operate.

The α -helix of a zinc finger peptide aligns antiparallel to the target nucleic acid strand, such that the primary nucleic acid sequence is arranged 3' to 5' in order to correspond 15 with the N-terminal to C-terminal sequence of the zinc finger peptide. Since nucleic acid sequences are conventionally written 5' to 3', and amino acid sequences N-terminus to C-terminus, the result is that when a target nucleic acid sequence and a zinc finger peptide are aligned according to convention, the primary interaction of the zinc finger peptide is with the “minus” strand of the nucleic acid sequence, since it is this strand 20 which is aligned 3' to 5'. These conventions are followed in the nomenclature used herein. It should be noted, however, that in nature certain zinc finger modules, such as zinc finger 4 of the protein GLI, bind to the “plus” strand of the nucleic acid sequence. See Suzuki *et al.* (1994) *Nucl. Acids Rev.* 22: 3397-3405; and Pavletich & Pabo, (1993) *Science* 261: 1701-1707. The present invention encompasses incorporation of such zinc 25 finger peptides into DNA binding molecules.

Natural Zinc Finger Peptides.

In certain embodiments, this invention relates to natural zinc finger modules. As used 30 herein, the term ‘natural’ with reference to a zinc finger, means that the DNA sequence which encodes a particular zinc finger, whether normally expressed *in vivo* or not, is found in nature, *i.e.* is part of the genome of a cell. A natural human zinc finger is one

which is endogenous to the human genome, a natural mouse zinc finger is found in the mouse genome, and a natural viral zinc finger is found in a viral genome, *etc.* Natural zinc finger genes which have become integrated into the genome of a heterologous species by natural means, *e.g.*, integration of a viral genome into a host genome, are

5 considered to be endogenous to the host species within the context of this disclosure. A zinc finger module constructed or produced *in vitro* or extracted from an *in vivo* source is considered to be natural if its amino acid sequence matches that of the amino acid sequence encoded by its natural gene. The DNA sequence of the natural gene is not the defining aspect. Thus, polynucleotides encoding natural zinc finger modules may have a
10 different sequence from that of the naturally-occurring sequence encoding the module, *e.g.*, to adjust codon usage to optimise expression of the module in a particular expression system.

15 Preferably, sequences of zinc fingers used in the present invention are not mutated from their natural form. Advantageously, the natural zinc finger polypeptides are expressed in nature.

A natural zinc finger binding motif is a structure well known to those in the art and defined in, for example, Miller *et al.*, (1985) *EMBO J.* 4: 1609-1614; Berg (1988) *Proc. 20 Natl. Acad. Sci. USA* 85: 99-102; Lee *et al.*, (1989) *Science* 245: 635-637; see also International patent applications WO 96/06166 and WO 96/32475, incorporated herein by reference.

In general, a natural zinc finger framework has the structure:

25 SEQ ID NO:12 X₀₋₂ C X₁₋₅ C X₉₋₁₄ H X₃₋₆ ^H/c

where X is any amino acid, and the numbers in subscript indicate the possible numbers of residues represented by X (Formula A).

In a preferred aspect of the present invention, natural zinc finger nucleic acid binding 30 motifs may be represented as motifs having the following primary structure:

X₀₋₂ C X₁₋₅ C X₂₋₇ X X X X X X H X₃₋₆ ^H/c (SEQ ID NO:14)
(SEQ ID NO:13)

- 1 1 2 3 4 5 6 7

where X is any amino acid, and the numbers in subscript indicate the possible numbers of residues represented by X (Formula A'). The numbers -1 through 7 refer to amino acid position with respect to the beginning of the alpha-helical region of the zinc finger.

5 The Cys and His residues, which together co-ordinate the zinc metal atom, are marked in bold text and are usually invariant. However, all naturally-occurring zinc finger modules, even if they diverge from the above formula, are encompassed within the scope of this invention.

10 Zinc finger modules of formula A' are often arranged in tandem within a natural zinc finger polypeptide, such that a zinc finger containing protein may have 2, 3, 4, 5, 6, 7, 8, 9 or more individual zinc finger motifs. In such a protein, individual zinc fingers are joined to each other by a polypeptide sequence known as a linker. Generally, such a natural linker lacks secondary structure, although the amino acids within the linker may 15 form local interactions when the protein is bound to its target site. By 'linker sequence' is meant an amino acid sequence that links together adjacent zinc finger modules. For example, in a natural zinc finger protein, the linker sequence is the amino acid sequence which lies between the last residue of the α -helix in a zinc finger and the first residue of the β - sheet in the next zinc finger. The linker sequence therefore joins together two zinc 20 fingers. For the purposes of the present invention, the last amino acid of the α -helix in a zinc finger is considered to be the final zinc coordinating histidine (or cysteine) residue, while the first amino acid of the following finger is generally a tyrosine / phenylalanine or another hydrophobic residue. Since some natural zinc fingers do not start with a hydrophobic residue (see Appendices), the start of a finger is sometimes harder to define 25 from amino acid sequence (or indeed zinc finger structure), and so some flexibility must be allowed in this definition. Accordingly, in a natural zinc finger protein, threonine is often considered to be the first residue in the linker, and proline is the last residue of the linker. Thus, for example, in the natural Zif268 peptide the linker sequence is - TG(E/Q)(K/R)P- (SEQ ID NO:15). Although natural linkers can vary greatly in terms of 30 amino acid sequence and length, on the basis of sequence homology, the canonical

natural linker sequence is considered to be -TGEKP- (SEQ ID NO:3). Hence, the preferred linker sequence to join zinc finger modules of the present invention is -TGEKP-.

5 Additionally, a 'leader' peptide may be added to the N-terminal zinc finger of a poly-zinc finger peptide to aid its expression, without changing the sequence of the natural zinc finger module. Preferably, the leader peptide is MAEERP (SEQ ID NO:16) or MAERP (SEQ ID NO:17).

10 In general, naturally occurring zinc finger modules may be selected from those proteins for which the DNA binding specificity is already known. For example, these may be the proteins for which a crystal structure has been resolved: namely Zif268 (Elrod-Erickson *et al.* (1996) *Structure* 4: 1171-1180), GLI (Pavletich & Pabo (1993) *Science* 261: 1701-1707), Tramtrack (Fairall *et al.* (1993) *Nature* 366: 483-487) and YY1 (Houbaviy *et al.* (1996) *Proc. Natl. Acad. Sci. USA* 93: 13577-13582). Furthermore, the sequence specificity of many naturally-occurring zinc fingers and zinc finger proteins are known. In addition, this invention further provides for the determination of the binding specificity of natural zinc finger modules for use in the present invention. *See "Prediction of Binding Specificity," infra.*

15

20

Poly-Zinc Finger Peptides.

It is desirable that a 'designer' transcription factor for uses such as gene therapy and in transgenic organisms should have the ability to target virtually unique sites within any genome. For complex genomes such as in humans, an address of at least 16 bps is required to specify a potentially unique DNA sequence. Shorter DNA sequences have a significant probability of appearing several times in a genome, raising the possibility of obtaining undesirable non-specific gene targeting with a designed transcription factor targeted to such a shorter sequence. As individual zinc fingers only bind 3 to 4 nucleotides, it is therefore necessary to construct multi-finger polypeptides to target these longer sequences. A six-zinc finger peptide (with an 18 bp recognition sequence) could, in theory, be used for the specific recognition of a single target site and hence, the

specific regulation of a single gene within any genome. In addition, a significant increase in binding affinity might also be expected, compared to a protein with fewer fingers. In simple terms, if a three-finger peptide (with a 9 bp recognition sequence) binds DNA with nanomolar affinity, two tandemly linked three-finger peptides might be expected to bind 5 an 18 bp sequence with an affinity of 10^{-15} - 10^{-18} M. However, most previous attempts at producing high-affinity 6-finger peptides (poly-zinc finger peptides) based on fusions of two 3-finger domains have been unsuccessful in generating much of an improvement in affinity over 3-finger peptides. Liu, Q., Segal, D. J., Ghiara, J. B. & Barbas, C. F. III (1997) *Proc. Natl. Acad. Sci. USA* 94: 5525-5530; Kim, J-S. & Pabo, C. O. (1998) *Proc. 10 Natl. Acad. Sci. USA* 95: 2812-2817; Kamiuchi, T., Abe, E., Imanishi, M., Kaji, T., Nagaoka, M. & Sugiura, Y. (1998) *Biochemistry* 37: 13827-13834. To optimise both the affinity and specificity of 6-finger peptides, a fusion of three 2-finger domains has been shown to be advantageous. Moore, M., Klug, A. & Choo, Y. (2001) *Proc. Natl. Acad. Sci. USA* 98: 1437-1441; and WO 01/53480. Therefore, in one embodiment, 2-finger 15 units are linked to make poly-zinc finger nucleotide-binding domains. A pool of 4096 such 2-finger units, that recognise all possible 6 bp sequences ($4^6=4096$), represents an archive sufficient to rapidly create universal nucleic acid recognition, by simple linkage, in an "off-the-shelf" manner. See Moore *et al.*, *supra* and WO 01/53480.

20 Poly-zinc finger peptides according to this invention may be constructed containing 2, 3, 4, 5, 6 or more zinc finger modules. Such poly-zinc finger peptides may contain inter-finger linkers other than the canonical (TGEKP) linker sequence, as described, for example, in WO 01/53479; Moore, M., Choo, Y. & Klug, A. (2001) *Proc. Natl. Acad. Sci. USA* 98: 1432-1436; and Moore, M., Klug, A. & Choo, Y. (2001) *Proc. 25 Natl. Acad. Sci. USA* 98: 1437-1441. Briefly, linker sequences may be flexible or structured but, in general, will not form base-specific interactions with the target nucleotide sequence. A 'flexible' linker is defined as one which does not form a specific secondary structure in solution, whereas a 'structured' linker is defined as one that adopts a particular secondary structure in solution. Preferably, flexible linkers include the 30 sequences GGERP (SEQ ID NO:18), GSERP (SEQ ID NO:19), GGGGSERP (SEQ ID NO:20), GGGGSGGSERP (SEQ ID NO:21), GGGGSGGSGGSERP (SEQ ID NO:22),

GGGGSGGSGGSGGSGGSERP (SEQ ID NO:23). Preferably, the structured linker comprises an amino acid sequence that is not capable of specifically binding nucleic acid. More preferably, the structured linker comprises the amino acid sequence of TFIIIA finger IV. Alternatively, or in addition, the structured linker is derived from a zinc finger 5 by mutation of one or more of its base contacting residues to reduce or abolish nucleic acid binding activity of the zinc finger. The zinc finger may be finger 2 of wild type Zif268 mutated at positions -1, 2, 3 and/or 6.

10 In one embodiment, this invention provides for the construction and screening of poly-zinc finger peptides containing at least one natural zinc finger module.

In another embodiment, this invention provides for the construction and screening of poly-zinc finger peptides containing at least one natural zinc finger module, linked with the canonical linker sequence -TGEKP- (SEQ ID NO:3).

15 In one embodiment, methods for the construction and use of poly-zinc finger peptide comprising natural zinc finger modules are provided.

20 In another embodiment, methods for the construction and use of poly-zinc finger peptide comprising natural zinc finger modules, linked with the canonical linker sequence -TGEKP- (SEQ ID NO:3), are provided.

25 In a further embodiment, methods for the construction and use of poly-zinc finger peptides comprising at least one natural zinc finger module, containing either flexible or structured linkers (as described above and in WO 01/53480), are provided.

b. Advantages of Natural Zinc Finger Modules

30 Zinc finger modules are compact and stable structures of approximately 30 amino acids, which contain the full information required to bind a nucleic acid triplet or overlapping quadruplet. As such, they have proven to be extremely versatile scaffolds for engineering novel DNA-binding domains. *See*, for example, Rebar, E. J. & Pabo, C. O. (1994)

Science 263, 671-673; Jamieson, A. C., Kim, S.-H. & Wells, J. A. (1994) Biochemistry 33, 5689-5695; Choo, Y. & Klug, A. (1994) Proc. Natl. Acad. Sci. U.S.A. 91, 11163-11167; Choo, Y., Sanchez-Garcia, I. & Klug, A. (1994) Nature 372, 642-645; Wu, H., Yang, W.-P. & Barbas III, C. F. (1995) Proc. Natl. Acad. Sci. USA 92, 344-348;

5 Greisman, H. A. & Pabo, C. O. (1997) Science 275, 657-661; Isalan, M., Klug, A. & Choo, Y. (1998) Biochemistry 37, 12026-12033; Choo, Y. (1998) Nature Struct. Biol. 5, 264-265; Segal, D. J., Dreier, B., Beerli, R. R. & Barbas, C. F. (1999) Proc. Natl. Acad. Sci. USA 96, 2758-2763; Isalan, M. & Choo, Y. (2000) J Mol Biol 295, 471-477; and Beerli, R. R., Dreier, B., Barbas, C.F. (2000) Proc Natl Acad Sci U S A 97, 1495-500.

10 The resulting engineered zinc finger domains have increased our knowledge of sequence-specific DNA recognition, as well as provided a wide range of potential tools for medicine and biotechnology.

15 As a result of these and other studies on zinc finger engineering, it has been recognised that an individual zinc finger module does not necessarily recognise a simple nucleotide triplet, as was first thought; but instead, can bind to an overlapping quadruplet of double stranded DNA. *See*, for example, Isalan *et al.* (1997) Proc Natl Acad Sci U S A 94, 5617-5621; and WO98/53057). In this respect, zinc finger engineering strategies have been particularly important for deciphering the mechanism and specificity of these interactions.

20 With the recent completion of the human genome project and the rapidly advancing fields of transgenic animals and plants, thousands of uncharacterised (and characterised) genes have (and will) become valid targets for functional genomics and other such projects. Concomitantly, engineered zinc finger peptides (often as a component of "designer"

25 transcription factors) are emerging as one of the most universal and desirable ways of regulating the expression of specific genes within cells. *See*, for example, Choo, Y., Sanchez-Garcia, I. & Klug, A. (1994) Nature 372: 642-645; Beerli, R. R., Dreier, B. & Barbas, C. F. III (2000) Proc. Natl. Acad. Sci. USA 97: 1495-1500; Kim, J-S. & Pabo, C. O. (1998) Proc. Natl. Acad. Sci. USA 95: 2812-2817; Kang, J. S. & Kim, J-S. (2000) J. Biol. Chem. 275: 8742-8748; Zhang *et al.* (2000) J. Biol. Chem. 275: 33,850-33,860; Liu *et al.* (2001) J. Biol. Chem. 276: 11,323-11,334; Ren *et al.* (2002) Genes. Devel. 16: 27-32; and WO 00/41566.

Notwithstanding the remarkable progress in zinc finger engineering, there remain several issues that limit the use of engineered zinc fingers for such applications. Points of particular concern include the potential immunogenicity of non-natural zinc fingers, and 5 the 'fine-tuning' of particular aspects of the protein-DNA interactions to obtain optimal and specific zinc finger-nucleic acid contacts.

The present invention overcomes problems such as immunogenicity and optimal binding specificity, by exploiting the vast repertoire of naturally occurring zinc fingers to 10 construct targeted zinc finger proteins having novel specificities.

Immunogenicity

The main function of the immune system is to detect, and render harmless, foreign 15 particles which have invaded the body as a whole, or individual cells or organs. 'Foreign' in this context means non-host, i.e. a substance which has originated from a different species, or one which has originated as a result of a mutation al event (such as might generate a malignant cell). On encountering such an antigenic particle, either in solution or on the surface of an infected cell, the body's defences rapidly destroy/remove it by 20 complex pathways which involve the interaction of many members of the immune system. For a good overview of immunology see Roitt, *Essential Immunology*, Blackwell Science Ltd. and Roitt, I., Brostoff, J. & Male, D. *Immunology*, 4th Ed. Mosby. Hence, all biological therapeutic agents, such as peptides, nucleic acids, viruses, etc., risk eliciting 25 an immune response in the recipient. Particularly for cases in which repeated doses of a therapeutic agent are required, this response can be strong and potentially dangerous to the host organism.

The immune system functions through either innate or adaptive responses. The innate response is usually the body's first internal line of defence. Phagocytic cells recognise 30 and bind to foreign objects in extracellular environments. Once bound, the foreign object is internalised and destroyed. Foreign therapeutic agents such as peptides and nucleic acids, which are administered directly to the blood stream of the recipient, risk being

detected and possibly destroyed before they even reach their intended target. This response is one of primitive non-specific recognition of non-host agents, and does not adapt with time or exposure to the antigen.

5 Foreign therapeutic agents (or infectious agents such as bacteria and viruses), which evade the innate immune response and may have been successfully delivered to a particular cell have not necessarily avoided the host's immune system. Proteins that are expressed in cells are routinely degraded within lysosomes, and short peptide fragments, generally of between 6 and 9 amino acids, are transported to the cell surface and
10 presented to the host's immune system. This is the start of the host's second internal defence mechanism against invasion, the adaptive immune response. The proteins responsible for displaying such peptide fragments are known as major-histocompatibility complexes (MHC) proteins. Lymphocyte cells, known as T-lymphocytes, dock with the MHC proteins and scan the peptide fragments displayed. Contact of a T-lymphocyte with
15 a fragment specifically recognised as not belonging to the host organism initiates an immunological cascade which ultimately results in the host cell being destroyed or undergoing apoptosis. This mechanism is one of specific recognition, and once recognised as foreign, the antigen is 'remembered' so that any future invasions by the agent are dealt with more and more rapidly. B-cells are another type of lymphocyte that
20 recognise extracellular particles and then produce and release antibodies to help combat the agent.

25 To avoid potentially damaging the host organism and to ensure the successful delivery and action of a therapeutic peptide it is important to make it as much like a host protein as is reasonably possible. In the case of synthesised therapeutic antibodies for human use, a great deal of work has gone in to the 'humanisation' of antibodies produced by other animal species (See EP 0239400). In this invention we present a solution for the equivalent problem associated with zinc finger therapeutic peptides.

30 To some extent, prior art zinc finger engineering strategies have attempted to minimise the risk of eliciting immune responses by using an engineering scaffold that is compatible with (*i.e.* that originates from) the recipient, and by limiting the sizes of the varied regions

within the final product. For example, typical engineered zinc fingers utilize a scaffold such as the three-finger DNA-binding domain of Zif268 (containing approximately 100 amino acid residues). Because the amino acid sequence of Zif268 is completely conserved in a variety of species, including mice and humans, the scaffold is not itself 5 immunogenic in these species. However, in order to engineer new DNA-binding domains, stretches of approximately 7 amino acids must be varied within each zinc finger. These sequences of 7 amino acids represent modifications in positions -1, 1, 2, 3, 4, 5, and 6 of the α -helix of each finger. Although these engineered regions are considered to be relatively small, they are approximately the length of the peptide 10 fragments displayed on the surface of cells by MHC molecules. Hence, they may provide antigenic peptide fragments in several registers of the amino acid sequence, which may result in dangerous and/or undesirable immune responses in the host.

Accordingly, it is not known whether this type of engineering strategy will be entirely 15 sufficient to avoid all potential undesirable effects, or indeed whether it will create the most optimal framework for all zinc finger-nucleic acid interactions.

In addition to the zinc fingers themselves, it is also possible that inter-finger linker sequences could present potential immunological problems. Fortunately, natural zinc 20 finger proteins display strong conservation and homology in their linker sequence. A very large number of natural fingers are joined by the canonical linker peptide -TGEKP- (SEQ ID NO:3), located between the final zinc chelating residue (usually histidine) of the first finger, and the first residue of the second finger (usually a large hydrophobic residue such as tyrosine or phenylalanine, which begins the β -sheet). Hence, the use of the 25 canonical linker sequence -TGEKP- (SEQ ID NO:3), to join natural zinc finger modules in a non-natural order, will reduce the possibility of eliciting an undesirable immune reaction to a minimum. Furthermore, there are so many natural zinc fingers which are already joined by canonical linker sequences, that if deemed necessary, the database of natural zinc fingers used for the construction of poly-zinc finger peptides may be 30 restricted to those already flanked by such linkers.

The periodicity of zinc fingers and their amenability to linkage using the TGEKP (SEQ ID NO:3) motif is illustrated in Table 2.

	α -HELIX	LINKER
5	-1123456	
	YA CPVESCDRRFS (SEQ ID NO:24) RSDELTR <u>H</u> IRIH (SEQ ID NO:25) <u>TGEKP</u>	
	FQ CRI CMRNFS (SEQ ID NO:26) RSDHLST <u>H</u> IRTH (SEQ ID NO:27) <u>TGEKP</u>	
	FA CDI CGRKFA (SEQ ID NO:28) RSDERKR <u>H</u> TKIH (SEQ ID NO:29) <u>TGEKP</u>	

10 **Table 2.** A functional three-finger DNA-binding domain based on the peptide sequence of Zif268. TGEKP linker motifs are underlined. The helical residues of each zinc finger are numbered relative to the first helical position, position +1. Conserved Cysteines and Histidines forming the classical Cys₂His₂ zinc finger core are shown in bold.

15

Fine-Tuning of Zinc Finger-Nucleic Acid Interactions.

It has previously been shown that zinc fingers cannot simply be regarded as independent nucleic acid-binding modules. Isalan, M., Klug, A. & Choo, Y. (1998) Biochemistry 37, 20 12026-12033; Isalan, M., Choo, Y. & Klug, A. (1997) Proc Natl Acad Sci 94, 5617-5621. The interactions between adjacent zinc fingers can be complex and involve overlap of binding sites, which means that optimal interfaces are not easily engineered through rational design. Combinatorial library selection systems, which if designed correctly necessarily result in interface compatibility, can help to engineer better optimisation of 25 the zinc finger-nucleic acid interface. *See*, for example, WO98/53057. However, all library selection systems suffer from the problem of library size, whereby because of physical constraints, it is impossible to include an exhaustive combination of randomisations to cover all potentially important sequence-space. For example, to optimise the zinc finger-nucleic acid interface, subtle amino acid variations may be 30 needed, even from positions outside the recognition α -helix. Furthermore, alternative approaches to zinc finger engineering, such as 'affinity maturation' through random mutation or gene shuffling, which may (to a limited extent) increase the coverage of sequence space, may also raise the probability of generating undesirable immunological problems. Hence, it is possible that the creation of truly optimal zinc finger domains for

recognition of specific nucleic acid sequences may be outside the scope of traditional engineering strategies.

In contrast, naturally occurring zinc finger modules have already been 'fine-tuned' by
5 thousands of years of natural selection and are, under normal circumstances, non-immunogenic in their host organism. The human genome project has revealed that zinc finger-containing proteins constitute the second most abundant family of proteins in humans, with well over 600 members. Since zinc finger proteins usually contain several individual zinc finger modules, the human genome provides a repertoire of thousands of
10 natural zinc finger modules for the creation of composite binding polypeptides.

Furthermore, because there are only 64 ($=4^3$) possible 3 bp sequences and 256 ($=4^4$) possible 4 bp sequences, it is likely that a natural zinc finger domain exists which is capable of binding to every potential 3- or 4-nucleotide target sequence. Consequently, natural zinc fingers are a very useful resource for the production of composite binding
15 polypeptides comprising zinc fingers. At present, the natural binding site of many natural zinc finger modules is not known. Thus, to be useful for the construction of composite binding polypeptides, nucleotide sequence preferences for certain natural zinc fingers are determined according to rules tables disclosed in the following section ("Binding Specificity of Natural Zinc Finger Modules").

20

To create optimal poly-zinc finger peptides the potentially significant problem of interface incompatibility must be addressed, since natural zinc finger modules will not necessarily be compatible with each other when juxtaposed. In this respect, a library construction and screening system is preferably employed which links natural zinc finger modules in non-natural combinations, and screens them against possible target sequences of greater than 3 or 4 bp in length (which represents the possible binding site of a single zinc finger module), to determine optimal 2- or 3-finger domains. In this way, the cooperative nature of zinc finger binding is taken into account in the design and selection of composite binding polypeptides, and in the determination of the sequence specificity of
25 their binding. In one embodiment, a library of poly-zinc finger peptides containing at least one natural zinc finger module is provided. Preferably, poly-zinc finger peptides of
30 the library contain at least two natural zinc finger modules.

5 c. **Binding Specificity of Natural Zinc Finger Modules**

Disclosed herein are certain improvements to current limitations on the use of customised zinc finger nucleic acid binding domains, through the use of natural zinc finger modules. By using either natural 1-finger or 2-finger sub-domains, and/or novel combinatorially-mixed, pre-selected 2-finger sub-domains, it is possible to construct poly-zinc finger peptides that bind any desired nucleotide target sequence, using non-natural combinations of natural zinc fingers.

This approach is particularly suited for human gene therapy applications, but the invention is not just limited to zinc finger modules encoded by the human genome. For applications within transgenic animals such as mice, chicken, etc., the same system can be used, but incorporating natural zinc finger modules from those species instead (see Example 3). The genome of any organism (e.g., animal, plant, bacterium, virus, etc.) can thus provide a genetic 'toolbox' of non-immunogenic, structurally optimised zinc fingers for applications in that organism.

Before such zinc finger modules can be utilised, however, it is essential that their optimal binding site is determined, in isolation, or preferably as part of a 2- or 3-finger subdomain. Natural zinc finger modules are advantageously fused into subdomains comprising two or three zinc finger modules in random arrangement, optionally comprising an anchor finger, then subjected to binding site analysis. An 'anchor' zinc finger is one for which the binding specificity is known, such as, for example, finger 1 or finger 3 of Zif268, each of which binds the sequence 5'-GCG-3'. An anchor finger is attached to the N- or C-terminus of the zinc finger module(s) or subdomain for which the binding specificity is to be determined, and acts as an anchor to set the binding register for the binding site selection. For example, if the binding site preference of a pair of natural zinc fingers is to be determined, finger 1 of Zif268 may be fused to the N-

terminus of the pair of natural fingers, and a 5'-GCG-3' anchor sequence is placed at the 3' end of 6 or more randomised nucleotides. Selection of the optimal binding site may thus be conducted with an oligonucleotide containing the sequence 5'-XXX-XXX-GCG-3' (SEQ ID NO:30), where X is any specified nucleotide. The anchor sequence thereby 5 allows the binding site preference of the zinc finger libraries to be easily determined. Such procedures are described in the Examples.

Screening for Zinc Finger Binding Specificity

10 There are various approaches, known to those in the art, for screening nucleic acid binding peptides for their binding specificity. To determine the binding specificity of, for example, zinc finger peptides, procedures can be conducted using: (a) a library of zinc fingers and a specified target sequence – to select one or more zinc finger peptides with a particular binding preference; or (b) a single zinc finger peptide and a random population 15 of target sequences – to select one or more optimal binding sites for a particular peptide. For many applications, such as for the creation of transcription factors for regulating specific gene activity, it is often preferable to screen zinc finger libraries against specific target sequences. In this way, the search is geared towards a particular application. However, if the function or binding specificity of a natural protein is the object of the 20 investigation, a library of potential binding sites can be screened using a single peptide. Some such methods are outlined below.

A typical method for screening libraries of nucleic acid binding polypeptides against specific target sites is that of phage display. Phage display protocols generally involve 25 expressing the peptides under study as fusions with the gIII major coat protein of bacteriophage (J. McCafferty, R. H. Jackson, D. J. Chiswell, (1991) *Protein Engineering* 4, 955-961). Suitable protocols for the selection of zinc finger peptides have been described and are well known to those in the art. *See*, for example, Choo, Y. & Klug, A. (1994) Proc. Natl. Acad. Sci. U.S.A. 91, 11163-11167; Choo, Y., Sanchez-Garcia, I. & 30 Klug, A. (1994) Nature 372, 642-645; Choo, Y. (1998) Nature Struct. Biol. 5, 264-265; Choo, Y. & Klug, A. (1997) Curr. Opin. Str. Biol. 7, 117-125; Isalan, M., Klug, A. & Choo, Y. (1998) Biochemistry 37, 12026-12033; Isalan, M. & Choo, Y. (2000) J Mol

Biol 295, 471-477; Isalan, M., Choo, Y. & Klug, A. (1997) Proc Natl Acad Sci 94, 5617-5621; WO 01/53480, WO 01/53479, WO 96/06166, WO 98/53057, WO 98/53058, WO 98/53059 and WO 98/53060 and references cited therein; see also Examples, *infra*. In general, sequences comprising target sites are bound, such as through biotin-streptavidin, 5 to a solid support, such as a magnetic particle, or the surface of a tube or well. A solution of phage expressing members of a library of zinc finger peptides is then added to the immobilised target site. Non-bound phage are washed away and bound phage (containing the DNA encoding the bound zinc finger peptide), are collected. The collected phage sample is usually reused in further rounds of selection to enrich for the tightest binding 10 zinc finger peptide.

Phage display protocols based on random mutagenesis of zinc finger modules are known to have a number of limitations. First, as discussed above, the library size that can be expressed on the surface of phage is limited by the efficiency of procedures such as 15 cloning and transformation. Furthermore, the efficiency of incorporation of gIII-zinc finger fusions into phage and hence, zinc finger peptide expression, is determined by the number of zinc finger modules. Therefore, 2-finger peptides are expressed more efficiently than 3-finger peptides and so on. For this reason, phage display protocols are generally limited to the assay of polypeptides comprising 3 or fewer zinc finger modules.

20 An alternative to phage display is an *in vitro* selection system. In such a system, libraries of zinc fingers can be produced by PCR using degenerate primer oligonucleotides. Target binding sites are added to the end of the DNA encoding the zinc finger peptide. Zinc finger peptide expression may be performed directly from PCR products using an *in* 25 *vitro* expression kit, such as the TNT T7 Quick Coupled Transcription/Translation System for PCR DNA (Promega, Madison, WI, USA), or another suitable expression system. The components of the expression reaction (including the zinc finger gene/binding site) are compartmentalised by suspension in an emulsion, in such a way that (on average) only one copy of the zinc finger gene / binding site is present in each 30 compartment. *See*, for example, Tawfik, D.S. & Griffiths, A.D. (1998) Nat. Biotechnol. 16: 652-656. Zinc finger peptides which bind the specified target site (and the gene encoding them) can be collected using, for example, a suitable epitope tag (such as myc,

FLAG or HA tags), and the non-bound binding sites/zinc finger genes are removed. The genes encoding zinc finger peptides that bind the required target site can then be amplified by PCR and used in further rounds of selection if required.

5 A preferred method for selecting a zinc finger peptide which binds a specified target sequence is described in Example 4. Briefly, the DNA encoding a library of zinc finger peptides with an attached epitope tag is diluted into as many aliquots as it is possible to screen (e.g. 384 or 1534 aliquots). This creates pools of sub-libraries with reduced numbers of variants. The DNA is then amplified by PCR and used to produce protein,

10 from a suitable *in vitro* expression system, as described above. A specified binding site with an attached biotin molecule, and a horse radish peroxidase (HRP)-conjugated antibody to the peptide-attached epitope tag may then be added. Binding site / bound zinc finger / antibody complexes may be collected by binding to streptavidin and the samples are washed to remove unbound zinc finger and antibodies. The samples

15 containing the highest amount of bound zinc finger peptide can be detected by adding an HRP substrate solution. The original DNA stock from such positive samples may then be diluted into aliquots (as above), PCR-amplified and used for the next round of selection. In this way, pools of zinc finger encoding genes with the desired activity are isolated, subdivided into pools of reduced variation and re-isolated until the most active clone is

20 identified.

Principal advantages of the *in vitro* systems described above are: (a) there is virtually no limit to the library size which can be screened (up to 10^{12} different PCR products can easily be made); and (b) polypeptides comprising larger numbers of linked zinc finger

25 modules (e.g., 4, 5, 6, 7, or more) can be assayed. Another *in vitro* selection system which can be used is polysome/ribosome display. *See*, for example, Mattheakis, L.C., Bhatt, R.R. & Dower, W.J. (1994) *Proc. Natl. Acad. Sci. USA* 91: 9022-9026; and WO 00/27878.

30 Protocols for the reverse selection procedure, *i.e.* the selection of a particular binding site from a mixed population using a single nucleic acid binding polypeptide, include SELEX (systematic evolution of ligands by exponential enrichment) and microarray techniques.

The SELEX procedure has been well described. See, for example, Drolet, D.W., Jenison, R.D., Smith, D.E., Pratt, D. & Hicke, B.J. (1999) Comb. Chem. High Throughput Screen 2: 271-278; Burden, D.A. & Osheroff, N. (1999) J. Biol. Chem. 274: 5227-5235;

5 Shultzaberger, R.K. & Schneider, T.D. (1999) Nucleic Acids Res. 27: 882-887; Marozzi, A., Meneveri, R., Giacca, M., Gutierrez, M.I., Siccardi, A.G. & Ginelli, E. (1998) J. Biotechnol. 15: 117-128; and US Patents No. 5,270,163; 5,475,096; 5,595,877; 5,670,637; 5,696,249; 5,817,785 and 6,331,398. A single nucleic acid binding polypeptide is expressed, either *in vitro* or *in vivo*, and screened against a library of target

10 sequences. Nucleic acid binding polypeptides are collected (along with any bound target sites) using an epitope tag (as above) or another suitable procedure. Bound target sites are amplified by PCR and may be used in further rounds of selection, to enrich for the optimal binding site, or sequenced.

15 Microarray technology provides a method of screening a particular polypeptide or nucleic acid against thousands to millions of target sequences on a single slid support such as, for example, a glass or nitrocellulose slide. For example, the members of a library encoding polypeptides comprising 2 linked zinc fingers will bind a 6 bp recognition sequence. Hence, there are 4096 ($=4^6$) unique binding sites for such a library. All 4096 of these

20 sites can be arrayed onto a single glass slide, for example, allowing a specified 2-finger peptide to be screened simultaneously against every possible binding site. The amount of binding to each target sequence can be visualised and quantified using simple fluorescence measurements. For example, the zinc finger peptide may be expressed *in vitro*, or on the surface of phage. Isolated zinc finger peptides may contain an epitope tag

25 for labelling purposes, whereas bound phage can be detected using a primary antibody against a phage coat protein, such as gVIII. A secondary antibody conjugated to, for example, R-phycoerythrin, horseradish peroxidase or alkaline phosphatase, can be used to provide a visible, quantifiable signal when a suitable substrate is applied. See, for example, Bulyk *et al.* (2001) Proc. Natl. Acad. Sci. USA:98,:13, 7158-7163, which is

30 incorporated, by reference, in its entirety.

Prediction of Binding Specificity

The screening approaches described above rely on the assay of large libraries of randomly-selected natural zinc finger modules, to obtain one or more zinc finger modules 5 that optimally bind a particular target nucleic acid sequence. In order to simplify the process further and ensure a more rapid selection of optimal zinc finger modules for a particular target site, sub-libraries can be created. In this disclosure, the term 'sub-library' refers to a library of natural zinc finger modules that have been roughly categorised according to their predicted binding specificity. For example, the total 10 population of natural zinc fingers can be sub-divided to create libraries comprising zinc finger modules whose predicted binding sites are guanine (G) rich, cytosine (C) rich, adenine (A) rich or thymine (T) rich. Alternatively, sub-libraries can be categorised as binding G in the 3' position, in the central position, or in the 5' position of a nucleotide triplet, etc. Alternatively, sub-libraries can be created which comprise zinc finger 15 modules predicted to bind a particular triplet sequence such as, for example, GGG, GGA, GGC, GGT, GAG, GCG, GTG, etc. This approach combines knowledge of the modes of zinc finger-nucleic acid recognition, gained from studies on artificial zinc finger variants, with the benefits of combinatorial library selection. It also takes into account the fact that concerted interactions between adjacent zinc fingers, i.e. overlapping contacts, can affect 20 the binding affinity and/or specificity of individual zinc fingers. *See*, for example, Isalan, M., Klug, A. & Choo, Y. (1998) Biochemistry 37, 12026-12033; Isalan, M., Choo, Y. & Klug, A. (1997) Proc Natl Acad Sci 94, 5617-5621. Thus, for example, a composite binding polypeptide comprising two fingers, each having a predicted binding specificity for a particular triplet, can be easily screened to determine if that pair of 25 fingers are compatible with each other for binding to the 6-nucleotide target site comprising their individual target sequences. This strategy is described further in the Examples.

For the process of creating sub-libraries of natural zinc fingers according to predicted 30 binding preference, the rules set forth in international patent applications WO 96/06166, WO 98/53057, WO 98/53058, WO 98/53059 and WO 98/53060, and described in more detail below, are used. These rules allow the assignment of an amino acid residue, in an

appropriate position of the recognition region of a zinc finger module (generally comprising amino acids -1 through +6, with respect to the start of the alpha-helical portion of the finger), which will bind a specified nucleotide in a triplet or quadruplet target subsite. However, these rules can also be used to predict the sequence of a target

5 subsite that would be preferentially bound by a zinc finger of given amino acid sequence. In particular, the identity of the amino acid residing at a particular position in the recognition region of a natural zinc finger module can be used to predict the identity of a nucleotide at a particular location in a target subsite. These 'rules' should be considered as a guide to target site preference and not a guaranteed prediction, as binding site

10 specificity may be determined by variations elsewhere in the zinc finger module (i.e. outside of the recognition region), may be influenced by context, or may be influenced by factors as yet unknown. It should also be noted that some rules may be more generally applicable than others.

15 In the application of these rules, it should be noted that the recognition region of a zinc finger aligns such that the N-terminal to C-terminal sequence of the finger is arranged along the nucleic acid strand to which it binds in a 3'-to-5' direction. As a result, when a zinc finger sequence and a nucleic acid sequence (to which the finger binds) are aligned, the primary interactions occur between the zinc finger and the 'minus' strand of the

20 nucleic acid sequence (i.e. the strand which has a 3'-to-5' orientation). Furthermore, as stated above, the recognition region of a zinc finger comprises amino acids -1 through +6, with respect to the start of the alpha-helical portion of the finger. With respect to a particular zinc finger, an amino acid residue designated ++2 refers to the residue present in the adjacent (in the C-terminal direction) zinc finger, which (in certain instances)

25 buttresses an amino acid-nucleotide interaction and/or participates in a cross-strand interaction with a nucleotide.

Thus, the following set of rules can be used to predict a 3 bp target subsite for a given natural zinc finger module: (a) if the 5' base in the triplet is G, then position +6 in the α -helix is Arg; or position +6 is Ser or Thr and position ++2 is Asp; (b) if the 5' base in the triplet is A, then position +6 in the α -helix is Gln and ++2 is not Asp; (c) if the 5' base in the triplet is T, then position +6 in the α -helix is Ser or Thr and position ++2 is Asp; (d) if

the 5' base in the triplet is C, then position +6 in the α -helix may be any amino acid, provided that position +2 in the α -helix is not Asp; (e) if the central base in the triplet is G, then position +3 in the α -helix is His; (f) if the central base in the triplet is A, then position +3 in the α -helix is Asn; (g) if the central base in the triplet is T, then position +3 in the α -helix is Ala, Ser or Val; provided that if it is Ala, then one of the residues at -1 or +6 is a small residue; (h) if the central base in the triplet is C, then position +3 in the α -helix is Ser, Asp, Glu, Leu, Thr or Val; (i) if the 3' base in the triplet is G, then position -1 in the α -helix is Arg; (j) if the 3' base in the triplet is A, then position -1 in the α -helix is Gln; (k) if the 3' base in the triplet is T, then position -1 in the α -helix is Asn or Gln; (l) if the 3' base in the triplet is C, then position -1 in the α -helix is Asp.

Furthermore, a natural zinc finger module may be capable of binding specifically to a four-nucleotide target subsite that overlaps with the target subsite of an adjacent zinc finger. In this case a different set of 'rules' can be used to determine predicted binding sites for each zinc finger module. Accordingly, in the description below, the overlapping 15 4 bp binding site is described such that position 4 is the 5' base of a typical triplet binding site, position 3 is the central position of a typical triplet, position 2 is the 3' position of a typical triplet, and position 1 is the complement of the nucleotide which is contacted by the cross strand interaction from the +2 position of the zinc finger module. Position 1 can also be considered to be the 5' base of the triplet or quadruplet contacted by an adjacent 20 (in the N-terminal direction) finger, if present.

Binding to each base of a quadruplet by an α -helical zinc finger nucleic acid binding motif in a natural protein can be predicted with reference to the following rules: (a) if base 4 in the quadruplet is G, then position +6 in the α -helix is Arg or Lys; (b) if base 4 in the quadruplet is A, then position +6 in the α -helix is Glu, Asn or Val; (c) if base 4 in the 25 quadruplet is T, then position +6 in the α -helix is Ser, Thr, Val or Lys; (d) if base 4 in the quadruplet is C, then position +6 in the α -helix is Ser, Thr, Val, Ala, Glu or Asn; (e) if base 3 in the quadruplet is G, then position +3 in the α -helix is His; (f) if base 3 in the quadruplet is A, then position +3 in the α -helix is Asn; (g) if base 3 in the quadruplet is T, then position +3 in the α -helix is Ala, Ser or Val; provided that if it is Ala, then one of the

residues at -1 or +6 is a small residue; (h) if base 3 in the quadruplet is C, then position +3 in the α -helix is Ser, Asp, Glu, Leu, Thr or Val; (i) if base 2 in the quadruplet is G, then position -1 in the α -helix is Arg; (j) if base 2 in the quadruplet is A, then position -1 in the α -helix is Gln; (k) if base 2 in the quadruplet is T, then position -1 in the α -helix is

5 His or Thr; (l) if base 2 in the quadruplet is C, then position -1 in the α -helix is Asp or His; (m) if base 1 in the quadruplet is G, then position +2 is Glu; (n) if base 1 in the quadruplet is A, then position +2 Arg or Gln; (o) if base 1 in the quadruplet is C, then position +2 is Asn, Gln, Arg, His or Lys; (p) if base 1 in the quadruplet is T, then position +2 is Ser or Thr.

10 The above rules may be further refined to those described below: (a) if base 4 in the quadruplet is G, then position +6 in the α -helix is Arg; or position +6 is Ser or Thr and position ++2 is Asp; (b) if base 4 in the quadruplet is A, then position +6 in the α -helix is Gln and ++2 is not Asp; (c) if base 4 in the quadruplet is T, then position +6 in the α -helix is Ser or Thr and position ++2 is Asp; (d) if base 4 in the quadruplet is C, then 15 position +6 in the α -helix may be any amino acid, provided that position ++2 in the α -helix is not Asp; (e) if base 3 in the quadruplet is G, then position +3 in the α -helix is His; (f) if base 3 in the quadruplet is A, then position +3 in the α -helix is Asn; (g) if base 3 in the quadruplet is T, then position +3 in the α -helix is Ala, Ser or Val; provided that if it is Ala, then one of the residues at -1 or +6 is a small residue; (h) if base 3 in the quadruplet 20 is C, then position +3 in the α -helix is Ser, Asp, Glu, Leu, Thr or Val; (i) if base 2 in the quadruplet is G, then position -1 in the α -helix is Arg; (j) if base 2 in the quadruplet is A, then position -1 in the α -helix is Gln; (k) if base 2 in the quadruplet is T, then position -1 in the α -helix is Asn or Gln; (l) if base 2 in the quadruplet is C, then position -1 in the α -helix is Asp; (m) if base 1 in the quadruplet is G, then position +2 is Asp; (n) if base 1 in 25 the quadruplet is A, then position +2 is not Asp; (o) if base 1 in the quadruplet is C, then position +2 is not Asp; (p) if base 1 in the quadruplet is T, then position +2 is Ser or Thr.

The rules therefore predict that the presence of an Asp (D) residue at position +2 will preclude binding to either A or C by an amino acid at position +6 in an adjacent N-

30 terminal finger. Isalan, M., Klug, A. & Choo, Y. (1998) Biochemistry 37, 12026-12033;

Isalan, M., Choo, Y. & Klug, A. (1997) *Proc Natl Acad Sci* 94, 5617-56212. Therefore, natural zinc fingers containing Asp, Glu, Asn or Gln at +6 are likely to be incompatible with any C-terminal finger containing an Asp residue at position +2. Although there are many such rules to describe the overlap between adjacent zinc fingers, a certain degree of 5 degeneracy exists in these rules. Nonetheless, physical selection procedures (e.g., library construction and screening) can be used to extract optimal pairs of fingers for any given target subsite interface.

Not all natural zinc fingers have a DNA-binding function. For example, it is known that 10 many zinc fingers, such as those from TFIIIA, bind to RNA (Clemens, K. R. *et al.*, (1993) *Science* 260: 530-533; Bogenhagen, D.F. (1993) *Mol. Cell. Biol.* 13: 5149-5158; Searles, M. A. *et al.*, *J. Mol. Biol.* 301: 47-60 (2000)). The rules governing RNA binding by zinc fingers are less well understood than those of DNA binding, but some RNA binding zinc fingers can be identified on the basis of a characteristic sequence motif. Clemens, K. R. 15 *et al.*, (1993) *Science* 260: 530-533; Bogenhagen, D.F. (1993) *Mol. Cell. Biol.* 13: 5149-5158; Searles, M. A. *et al.* (2000) *J. Mol. Biol.* 301: 47-60. Furthermore, some zinc fingers, such as those from the protein Ikaros, are able to form protein-protein interactions. Such zinc fingers often contain large hydrophobic patches. Mackay, J. P. & Crossley, M. (1998) *Trends Biochem. Sci.* 23: 1-4.

20 To this end, applied bioinformatic processing can help to determine which candidates in a particular genome are best suited to fulfilling a particular function, such as DNA-binding. In the case of zinc fingers, numerous documented databases exist denoting amino acid residues that are most likely to be found at particular positions within a DNA-binding 25 zinc finger. See, for example, Isalan, M., Klug, A. & Choo, Y. (1998) *Biochemistry* 37, 12026-12033; Choo, Y. & Klug, A. (1997) *Curr. Opin. Str. Biol.* 7, 117-125; WO 98/53060; WO 98/53059; WO 98/53058. As an example, disclosed herein is a database of approximately 200 natural human zinc fingers which have been selected (on the basis of coded contacts) as having potentially useful DNA-binding activity (see Example 1). 30 Also disclosed in Example 1 are the predicted DNA target sequences of these zinc fingers, assigned according to the rules set out above.

As the human genome contains almost 700 zinc finger-containing proteins, there are many other candidates that can be included in a more inclusive library of natural zinc fingers. A selection of these are disclosed in Example 2.

5 Similar work can be carried out in other organisms, such as farm (cows, pigs, sheep, chickens, etc.), laboratory (monkeys, rats, mice, etc.) and domestic (dogs, cats, etc.) animals. In this case, it is necessary to select natural zinc finger modules from the respective genomes of such organisms. Examples of zinc finger modules which have been selected from mouse, chicken and certain plant genomes, are disclosed in

10 Example 3.

d. Zinc Finger Chimeric Peptides

In a preferred embodiment, the composite binding polypeptides described herein comprise chimeric nucleic acid binding polypeptides.

15 A chimeric nucleic acid binding polypeptide, also referred to as a fusion polypeptide, comprises a binding domain (comprising a number of nucleic acid binding polypeptide modules or fingers) designed to bind specifically to a target nucleotide sequence, together with one or more further biological effector domains or functional domains. The terms "biological effector domain" and "functional domain" refer to any polypeptide (of functional fragment thereof) that has a biological function. Included are enzymes, receptors, regulatory domains, transcriptional activation or repression domains, binding sequences, dimerisation, trimerisation or multimerisation sequences, sequences involved in protein transport, localisation sequences such as subcellular localisation sequences, nuclear localisation, protein targeting or signal sequences. Furthermore, 20 biological effector domains may comprise polypeptides involved in chromatin remodelling, chromatin condensation or decondensation, DNA replication, transcription, translation, protein synthesis, etc. Fragments of such polypeptides comprising the relevant activity (*i.e.*, functional fragments) are also included in this definition. Preferred biological effector domains include transcriptional modulation domains such as

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transcriptional activators and transcriptional repressors, as well as their functional fragments.

The effector domain(s) can be covalently or non-covalently attached to the binding domain.

5 Chimeric nucleic acid binding polypeptides preferably comprise transcription factor activity, for example, a transcriptional modulation activity such as transcriptional activation or transcriptional repression activity. For example, a zinc finger chimeric polypeptide may comprise a binding domain designed to bind specifically to a particular nucleotide sequence, and one or more further biological effector domains, preferably a
10 transcriptional activation or repression domain, as described in further detail below. The zinc finger chimeric polypeptide may comprise one or more zinc fingers or zinc finger binding modules.

15 Preferably, in the case of a chimeric polypeptide comprising transcriptional modulation activity, a nuclear localisation domain is attached to the DNA binding domain to direct the chimeric polypeptide to the nucleus.

Generally, a chimeric nucleic acid binding polypeptide, such as a chimeric zinc finger polypeptide, can also include an effector domain to regulate gene expression. The effector domain can be directly derived from a basal or regulated transcription factor such as, for example, transactivators, repressors, and proteins that bind to insulator or silencer sequences. See, for example, Choo & Klug (1995) *Curr. Opin. Biotech.* 6: 431-436; Choo, Y. & Klug, A. (1997) *Curr. Opin. Str. Biol.* 7, 117-125; Rebar & Pabo (1994) *Science* 263: 671-673; Jamieson *et al.* (1994) *Biochem.* 33: 5689-5695; Goodrich *et al* (1996) *Cell* 84: 825-830; Vostrov, A. A. & Quitschke, W. W. (1997) *J. Biol. Chem.* 272: 33353-33359 and WO 00/41566 and references disclosed therein. Other useful domains 20 are derived from receptors such as, for example, nuclear hormone receptors (Kumar, R & Thompson, E. B. (1999) *Steroids* 64: 310-319), and their co-activators and co-repressors 25 (Ugai, H. *et al.* (1999) *J. Mol. Med.* 77: 481-494).

A chimeric nucleic acid binding polypeptide can also include other domains that may be advantageous within the context of the control of gene expression. Such domains include, but are not limited to, protein-modifying domains such as histone acetyltransferases, kinases, methylases and phosphatases, which can silence or activate genes by modifying DNA structure or the proteins that associate with nucleic acids. *See*, for example, Wolffe, *Science* 272: 371-372 (1996); Taunton *et al.*, *Science* 272: 408-411 (1996); Hassig *et al.*, *Proc. Natl. Acad. Sci. USA* 95: 3519-3524 (1998); Wang, *Trends Biochem. Sci.* 19: 373-376 (1994); and Schonthal & Semin, *Cancer Biol.* 6: 239-248 (1995). Additional useful effector domains include those that modify or rearrange nucleic acid molecules such as methyltransferases, endonucleases, ligases, recombinases etc. *See*, for example, Wood, *Ann. Rev. Biochem.* 65: 135-167 (1996); Sadowski, *FASEB J.* 7: 760-767 (1993); Cheng, *Curr. Opin. Struct. Biol.* 5: 4-10 (1995); Wu *et al.* (1995) *Proc. Natl. Acad. Sci. USA* 92:344-348; Nahon & Raveh, *Nucleic Acids Res* 1998 Mar 1;26(5):1233-9; Smith *et al.* *Nucleic Acids Res.* 1999 Jan 15;27(2):674-81; and Smith *et al.* (2000) *Nucleic Acids Res.* Sept 1; 28(17):3361-9. It will be appreciated that the biological effector domain portion of the chimeric polypeptide may itself also comprise such activities, without the need for further additional domains.

For the purpose of gene activation, zinc finger domains may be fused to the VP64 domain. *See*, for example, Seipel *et al.*, *EMBO J.* 11: 4961-4968 (1996). Other preferred transactivator domains include the herpes simplex virus (HSV) VP16 domain (Hagmann *et al.* (1997) *J. Virol.* 71: 5952-5962; Sadowski *et al.* (1988) *Nature* 335:563-564), transactivation domain 1 and/or domain 2 of the p65 subunit of nuclear factor- κ B (NF- κ B (Schmitz, M. L. *et al.* (1995) *J. Biol. Chem.* 270: 15576-15584). Other transcription factors are reviewed in, for example, Lekstrom-Himes J. & Xanthopoulos K. G. (C/EBP family) *J. Biol. Chem.* 273: 28545-28548 (1998); Bieker, J. J. *et al.*, (globin gene transcription factors) *Ann. N. Y. Acad. Sci.* 850: 64-69 (1998), and Parker, M. G. (estrogen receptors) *Biochem. Soc. Symp.* 63: 45-50 (1998).

Use of a transactivation domain from the estrogen receptor is disclosed in Metivier, R., Petit, FG., Valotaire, Y. & Pakdel, F. (2000) *Mol. Endocrinol.* 14: 1849-1871. Furthermore, activation domains from the globin transcription factors EKLF

(Pandya, K. Donze, D. & Townes T. (2001) *J. Biol. Chem.* 276: 8239-8243) may also be used, as well as a transactivation domain from FKLF (Asano, H. Li, XS.& Stamatoyannopoulos, G. (1999) *Mol. Cell. Biol.* 19: 3571-3579). C/EPB transactivation domains may also be employed in the methods described herein. The C/EBP epsilon activation domain is disclosed in Verbeek, W., Gombart, AF, Chumakov, AM, Muller, C, Friedman, AD, & Koeffler, HP (1999) *Blood* 15: 3327-3337. Kowenz-Leutz, E. & Leutz, A. (1999) *Mol. Cell.* 4: 735-743 disclose the use of the C/EBP tau activation domain, while the C/EBP alpha transactivation domain is disclosed in Tao, H., & Umek, RM. (1999) *DNA Cell Biol.* 18: 75-84.

10 It is known that zinc finger proteins may be fused to transcriptional repression domains such as the Kruppel-associated box (KRAB) domain to form powerful repressors. These domains are known to repress expression of a reporter gene even when bound to sites a few kilobase pairs upstream from the promoter of the gene (Margolin *et al.*, 1994, *Proc. Natl. Acad. Sci. USA* 91: 4509-4513). Hence, in certain embodiments, 15 the KRAB repressor domain from the human KOX-1 protein is used to repress gene activity (Moosmann *et al.*, *Biol. Chem.* 378: 669-677 (1997); Thiesen *et al.*, *New Biologist* 2: 363-374 (1990)). In additional embodiments, larger fragments of the KOX-1 protein comprising the KRAB domain, up to and including full-length KOX protein, are used as transcriptional repression domains. *See*, for example, Abrink *et al.* (2001) *Proc. Natl. Acad. Sci. USA* 98:1422-1426. Other preferred transcriptional repressor domains are known in the art and include, for example, the *engrailed* domain (Han *et al.*, *EMBO J.* 12: 2723-2733 (1993)), the *snag* domain (Grimes *et al.*, *Mol Cell. Biol.* 16: 6263-6272 (1996)) and the transcriptional repression domain of v-erbA (*e.g.*, Urnov *et al.* (2000) *EMBO J.* 19:4074-4090; Sap *et al.* (1989) *Nature* 340:242-244 and Ciana *et al.* (1999) *EMBO J.* 17:7382-7394).

30 Biological effector domains can be covalently or non-covalently linked to a binding domain. In one embodiment, a covalent linker comprises a flexible amino acid sequence; fusion polypeptides according to this embodiment comprise a nucleic acid binding domain fused, by an amino acid linker, to a biological effector domain. Alternatively, a covalent linker may comprise a synthetic, non-amino acid based,

chemical linker, for example, polyethylene glycol. Synthetic linkers are commercially available, and methods of chemical conjugation are known in the art. Covalent linkers may comprise flexible or structured linkers, as described above.

Non-covalent linkages between a nucleic acid binding domain and an effector domain can be formed using, for example, leucine zipper/coiled coil domains, or other naturally occurring or synthetic dimerisation domains. *See e.g.*, Luscher, B. & Larsson, L. G. *Oncogene* 18:2955-2966 (1999) and Gouldson, P. R. *et al.*, *Neuropsychopharmacology* 23: S60-S77 (2000).

The expression of composite binding polypeptides (for example, zinc finger polypeptides) can be controlled by tissue specific promoter sequences such as, for example, the *lck* promoter (thymocytes, Gu, H. *et al.*, *Science* 265: 103-106 (1994)); the human CD2 promoter (T-cells and thymocytes, Zhumabekov, T. *et al.*, *J. Immunological Methods* 185: 133-140 (1995)); the alpha A-crystallin promoter (eye lens, Lakso, M. *et al.*, *Proc. Natl. Acad. Sci.* 89: 6232-6236 (1992)); the alpha-calcium-calmodulin-dependent kinase II promoter (hippocampus and neocortex, Tsien, J. *et al.*, *Cell* 87: 1327-1338 (1996)); the whey acidic protein promoter (mammary gland, Wagner, K.-U. *et al.*, *Nucleic Acids Res.* 25: 4323-4330 (1997)); the aP2 enhancer/promoter (adipose tissue, Barlow C. *et al.*, *Nucleic Acids Res.* 25: 2543-2545 (1997)); the aquaporin-2 promoter (renal collecting duct, Nelson R. *et al.*, *Am. J. Physiol.* 275: C216-C226 (1998)); and the mouse myogenin promoter (skeletal muscle, Grieshammer, U. *et al.*, *Dev. Biol.* 197: 234-247 (1998)). The expression of such polypeptides can also be controlled by inducible systems, in particular, controlled by small molecule induction such as the tetracycline-controlled systems (tet-on and tet-off), the RU-486 or tamoxifen hormone analogue systems, or the radiation-inducible early growth response gene-1 (EGR1) promoter. These promoter constructs and inducible systems have the benefit of being able to provide organ-specific and/or inducible expression of target genes for use in applications such as gene therapy and transgenic animals.

e. **Vectors**

The nucleic acid encoding the nucleic acid binding polypeptide such as a zinc finger polypeptide can be incorporated into intermediate vectors and transformed into prokaryotic or eukaryotic cells for expression or DNA amplification.

5 As used herein, vector (or plasmid) preferably refers to discrete elements that are used to introduce heterologous nucleic acid into cells for either expression or replication thereof. The term "heterologous to the cell" means that the sequence does not naturally exist in the genome of the host cell but has been introduced into the cell. The term "introduced into" means that a procedure is performed on a cell, tissue, organ or organism
10 such that the gene encoding the nucleic acid binding polypeptide (for example, a zinc finger polypeptide) previously absent from the cell or cells is then present in the cell or cells. Alternatively, or in addition, the gene may be initially present in the cell or cells and subsequently altered by introduction of heterologous DNA. A heterologous sequence may include a modified sequence introduced at any chromosomal site, or which is not
15 integrated into a chromosome, or which is introduced by homologous recombination such that it is present in the genome in the same position as the native allele. Selection and use of such vectors are well within the skill of the person of ordinary skill in the art. Many vectors are available, and selection of an appropriate vector will depend on the intended use of the vector, i.e. whether it is to be used for DNA amplification or for nucleic acid
20 expression, the size of the DNA to be inserted into the vector, and the host cell to be transformed with the vector, etc. Another consideration is whether the vector is to remain episomal or integrate into the host genome. Suitable vectors may be of bacterial, viral, insect or mammalian origin. Intermediate vectors for storage or manipulation of the nucleic acid encoding the nucleic acid binding polypeptide, or for expression and
25 purification of the polypeptide are typically of prokaryotic origin. Most expression vectors are shuttle vectors, i.e. they are capable of replication in at least one class of organisms but can be transfected into another class of organisms for expression. For example, a vector is cloned in *E. coli* and then the same vector is transfected into yeast or mammalian cells even though it is not capable of replicating independently of the host
30 cell chromosome. DNA may also be replicated by insertion into the host genome. The

nucleic acid binding polypeptides such as zinc finger polypeptides described here are preferably inserted into a vector suitable for expression in mammalian cells.

Prokaryote, yeast and higher eukaryote cells may be used for replicating DNA and producing the nucleic acid binding protein. Suitable prokaryotes include eubacteria, such 5 as Gram-negative or Gram-positive organisms, such as *E. coli*, e.g. *E. coli* K-12 strains, DH5a and HB101, or Bacilli. Further hosts suitable for the vectors include eukaryotic microbes such as filamentous fungi or yeast, e.g. *Saccharomyces cerevisiae*. Higher eukaryotic cells include insect and vertebrate cells, particularly mammalian cells including human cells or nucleated cells from other multicellular organisms. In recent 10 years propagation of vertebrate cells in culture (tissue culture) has become a routine procedure. Examples of useful mammalian host cell lines are epithelial or fibroblastic cell lines such as Chinese hamster ovary (CHO) cells, NIH 3T3 cells, HeLa cells or 293T cells. The host cells referred to in this disclosure comprise cells in *in vitro* culture as well as cells that are within a host animal.

15 Each vector contains various components depending on its function (amplification of DNA or expression of DNA) and the host cell for which it is compatible. The vector components generally include, but are not limited to, one or more of the following: an origin of replication, one or more selectable marker genes, a promoter, an enhancer element, a transcription termination sequence and a signal sequence.

20 Both expression and cloning vectors generally contain nucleic acid sequence that enable the vector to replicate in one or more selected host cells. Typically in cloning vectors, this sequence is one that enables the vector to replicate independently of the host chromosomal DNA, and includes origins of replication or autonomously replicating sequences. Such sequences are well known for a variety of bacteria, yeast and viruses. 25 The origin of replication from the plasmid pBR322 is suitable for most Gram-negative bacteria, the 2 μ plasmid origin is suitable for yeast, and various viral origins (e.g. SV 40, polyoma, adenovirus) are useful for cloning vectors in mammalian cells. Generally, the origin of replication component is not needed for mammalian expression vectors unless

these are used in mammalian cells competent for high level DNA replication, such as COS cells.

Advantageously, an expression and cloning vector contains a selection gene also referred to as selectable marker. This gene encodes a protein necessary for the survival or 5 growth of transformed host cells grown in a selective culture medium. Host cells not transformed with the vector containing the selection gene will not survive in the culture medium. Typical selection genes encode proteins that confer resistance to antibiotics and other toxins, e.g. ampicillin, neomycin, methotrexate or tetracycline, complement auxotrophic deficiencies, or supply critical nutrients not available from complex media.

10

Since the replication of vectors is conveniently done in *E. coli*, an *E. coli* genetic marker and an *E. coli* origin of replication are advantageously included. These can be obtained from *E. coli* plasmids, such as pBR322, Bluescript[®] vector or a pUC plasmid, e.g. pUC18 or pUC19, which contain both *E. coli* replication origin and *E. coli* genetic 15 marker conferring resistance to antibiotics, such as ampicillin and tetracycline. Vectors such as these are commercially available.

20

As to a selective gene marker appropriate for yeast, any marker gene can be used which facilitates the selection for transformants due to the phenotypic expression of the 20 marker gene. Suitable markers for yeast are, for example, those conferring resistance to antibiotics G418, hygromycin or bleomycin, or provide for prototrophy in an auxotrophic yeast mutant, for example the URA3, LEU2, LYS2, TRP1, or HIS3 gene.

25

Suitable selectable markers for mammalian cells are those that enable the identification of cells competent to take up nucleic acid, such as dihydrofolate reductase (DHFR, methotrexate resistance), thymidine kinase, or genes conferring resistance to neomycin, G418 or hygromycin. The mammalian cell transformants are placed under selection pressure which only those transformants which have taken up and are expressing the marker are uniquely adapted to survive. In the case of a DHFR or 30 glutamine synthase (GS) marker, selection pressure can be imposed by culturing the transformants under conditions in which the pressure is progressively increased, thereby

leading to amplification (at its chromosomal integration site) of both the selection gene and the linked DNA that encodes the nucleic acid binding protein. Amplification is the process by which genes in greater demand (such as one encoding a protein that is critical for growth), together with closely associated genes (such as one encoding a composite binding polypeptide), are reiterated in tandem within the chromosomes of recombinant 5 cells. Increased quantities of desired protein are usually synthesised from this amplified DNA.

Expression and cloning vectors usually contain control sequences that are recognised by the host organism and are operably linked to the nucleic acid encoding a 10 nucleic acid binding polypeptide. The term "control sequences" is intended to include, at a minimum, components whose presence can influence expression, and can also include additional components whose presence is advantageous, for example, leader sequences and fusion partner sequences. The term "operably linked" means that the components described are in a relationship permitting them to function in their intended manner. 15 Typical control sequences include promoters, enhancers and other expression regulation signals such as terminators. Such a promoter may be inducible or constitutive. A regulatory sequence operably linked to a coding sequence is ligated in such a way that expression of the coding sequence is achieved under conditions compatible with the control sequences.

20 The term promoter is well known in the art and encompasses nucleic acid regions ranging in size and complexity from minimal promoters to promoters including upstream elements and enhancers. Suitable promoters for use in prokaryotic and eukaryotic cells are well known in the art, and described in for example, *Current Protocols in Molecular Biology* (Ausubel *et al.*, eds., 1994) and *Molecular Cloning. A Laboratory Manual* 25 (Sambrook *et al.*, 2nd ed. 1989).

Promoters suitable for use with prokaryotic hosts include, for example, the β -lactamase and lactose promoter systems, alkaline phosphatase, the tryptophan (Trp) promoter system and hybrid promoters such as the tac promoter. Their nucleotide sequences have been published, thereby enabling the skilled worker to ligate them to

DNA encoding a composite binding protein, using linkers or adapters to supply any required restriction sites. Promoters for use in bacterial systems will also generally contain an adjacent ribosome binding site (e.g., a Shine-Dalgarno sequence) operably linked to the DNA encoding the composite binding polypeptide.

5 Preferred expression vectors are bacterial expression vectors, which comprise a promoter of a bacteriophage such as phage lambda, SP6, T3 or T7, for example, which is capable of functioning in bacteria. In one of the most widely used expression systems, the nucleic acid encoding the fusion protein can be transcribed from a vector by T7 RNA polymerase (Studier *et al*, *Methods in Enzymol.* 185: 60-89, 1990). In the *E. coli*
10 BL21(DE3) host strain, used in conjunction with pET vectors, the T7 RNA polymerase is produced from the λ -lysogen DE3 in the host bacterium, and its expression is under the control of the IPTG inducible lac UV5 promoter. This system has been employed successfully for over-production of many proteins. Alternatively, the polymerase gene may be introduced on a lambda phage by infection with an *int*⁺ phage such as the CE6
15 phage, which is commercially available (Novagen, Madison, WI, USA). Other vectors include vectors containing the lambda P_L promoter such as PLEX (Invitrogen, NL), vectors containing the *trc* promoters such as pTrcHisXpressTm (Invitrogen), or pTrc99 (Pharmacia Biotech, SE), or vectors containing the *tac* promoter such as pKK223-3 (Pharmacia Biotech), or PMAL (New England Biolabs, Beverly, MA, USA). A suitable
20 vector for expression of proteins in mammalian cells is the CMV enhancer-based vector such as pEVRF (Matthias, *et al.*, (1989) *Nucleic Acids Res.* 17, 6418).

Suitable promoting sequences for use with yeast hosts may be regulated or constitutive and are preferably derived from a highly expressed yeast gene, especially a *Saccharomyces cerevisiae* gene. Thus, the promoter of the *TRP1* gene, the *ADH1* or
25 *ADH2* gene, the acid phosphatase (PH05) gene, a promoter of the yeast mating pheromone genes coding for the α - or α -factor or a promoter derived from a gene encoding a glycolytic enzyme such as the promoter of the enolase, glyceraldehyde-3-phosphate dehydrogenase (GAP), 3-phosphoglycerate kinase (PGK), hexokinase, pyruvate decarboxylase, phosphofructokinase, glucose-6-phosphate isomerase, 3-
30 phosphoglycerate mutase, pyruvate kinase, triose phosphate isomerase, phosphoglucose

isomerase or glucokinase genes, or a promoter from the TATA binding protein (TBP) gene can be used. Furthermore, it is possible to use hybrid promoters comprising upstream activation sequences (UAS) of one yeast gene and downstream promoter elements including a functional TATA box of another yeast gene, for example a hybrid 5 promoter including the UAS(s) of the yeast PH05 gene and downstream promoter elements including a functional TATA box of the yeast GAP gene (PH05-GAP hybrid promoter). A suitable constitutive PH05 promoter is, for example, a shortened acid phosphatase PH05 promoter devoid of the upstream regulatory elements (UAS) such as the PH05 (-173) promoter element starting at nucleotide -173 and ending at nucleotide -9 10 of the PH05 gene.

The promoter is typically selected from promoters which are found in animal cells, although prokaryotic promoters and promoters functional in other eukaryotic cells can be used. Typically, the promoter is derived from viral or animal gene sequences, may be constitutive or inducible, and may be strong or weak.

15 Viral promoters can be derived from viruses such as polyoma virus, adenoviruses, adeno-associated viruses, poxviruses (e.g., fowlpox virus), papilloma viruses (e.g., BPV), avian sarcoma virus, cytomegalovirus (CMV), herpesviruses, retroviruses, lentiviruses and simian virus 40 (SV40). An example of a relatively weak viral promoter is thymidine kinase promoter from herpes simplex virus (HSV-TK).

20 Mammalian derived promoters can be heterologous to the animal in which composite binding polypeptide (such as zinc finger polypeptide) expression is to occur, or they can be host sequences. In some applications it is preferable to use a promoter that is active in all cell types, however it is often preferable to use promoter sequences that are active in specific cell types only.

25 The actin promoter and the strong ribosomal protein promoter are examples of promoter sequences that are active in all cell types. In contrast, by using promoters that are specific for certain cell or tissue types, the gene encoding the nucleic acid binding polypeptide can be expressed only in the required cell or tissue types. This may be of

extreme importance for applications such as gene therapy, and for the production of viable transgenic animals. Such promoters are known in the art and include the *lck* promoter (thymocytes, Gu, H. *et al.*, *Science* 265: 103-106 (1994)), the human CD2 promoter (T-cells and thymocytes, Zhumabekov, T. *et al.*, *J. Immunological Methods* 185: 133-140 (1995)); the alpha A-crystallin promoter (eye lens, Lakso, M. *et al.*, *Proc. Natl. Acad. Sci.* 89: 6232-6236 (1992)), the alpha-calcium-calmodulin-dependent kinase II promoter (hippocampus and neocortex, Tsien, J. *et al.*, *Cell* 87: 1327-1338 (1996)), the whey acidic protein promoter (mammary gland, Wagner, K.-U. *et al.*, *Nucleic Acids Res.* 25: 4323-4330 (1997)), the aP2 enhancer/promoter (adipose tissue, Barlow C. *et al.*, *Nucleic Acids Res.* 25: 2543-2545 (1997)), the aquaporin-2 promoter (renal collecting duct, Nelson R. *et al.*, *Am. J. Physiol.* 275: C216-C226 (1998)), the mouse myogenin promoter (skeletal muscle, Grieshammer, U. *et al.*, *Dev. Biol.* 197: 234-247 (1998)), retinoblastoma gene promoter (nervous system, Jiang, Z. *et al.*, *J. Biol. Chem.* 276: 593-600 (2001)).

15 The expression of nucleic acid binding polypeptides such as zinc finger polypeptides can also be controlled by small molecule induction or other inducible systems such as the tetracycline inducible systems (tet-on and tet-off), the RU-486 or tamoxifen hormone analogue systems, or the radiation-inducible early growth response gene-1 (EGR1) promoter, all of which are commercially available. By using such 20 inducible promoter systems, transgenic lines can be established which carry a zinc finger chimeric polypeptide but express it only after addition of an inducer molecule. Thus the genes encoding the zinc finger polypeptides or other nucleic acid binding polypeptides can be expressed (or not expressed) in response to the small molecule, which can be easily administered. These systems may also allow the time and amount of polypeptide 25 expression to be regulated.

Expression vectors typically contain expression cassettes that carry all the additional elements required for efficient expression of the nucleic acid in the host cell. Additional elements are enhancer sequences, polyadenylation and transcriptional termination signals, ribosome binding sites, and translational termination sequences.

Transcription of DNA by higher eukaryotes may be increased by inserting an enhancer sequence into the vector. Enhancers are relatively orientation and position independent. Many enhancer sequences are known from mammalian genes (e.g. elastase and globin). However, typically one will employ an enhancer from a eukaryotic cell 5 virus. Examples include the SV40 enhancer on the late side of the replication origin (approx. bp 100-270) and the CMV early promoter enhancer. The enhancer may be spliced into the vector at a position 5' or 3' to the gene encoding the zinc finger polypeptide or nucleic acid binding polypeptide, but is preferably located at a site 5' from the promoter.

10 It has also been shown that the expression of a heterologous gene in an animal cell may be enhanced by retaining intron sequences (as opposed to using a cDNA clone). For example, intron 1 of the human CD2 gene has been shown to enhance the level of expression of CD2 in human cells (Festenstein, R. *et al.* 1996 *Science* 271: 1123).

15 Advantageously, a eukaryotic expression vector encoding a nucleic acid binding protein may comprise a locus control region (LCR). LCRs are capable of directing high-level integration site-independent expression of transgenes integrated into host cell chromatin. This is particularly important where the gene encoding the zinc finger polypeptide or the nucleic acid binding polypeptide is to be expressed over extended periods of time, for applications such as transgenic animals and gene therapy, as gene 20 silencing of integrated heterologous DNA – especially of viral origin – is known to occur (Palmer, T. D. *et al.*, *Proc. Natl. Acad. Sci. USA* 88: 1330-1334 (1991); Harpers, K. *et al.*, *Nature* 293: 540-542 (1981); Jahner, D. *et al.*, *Nature* 298: 623-628 (1992); and Chen, W. Y. *et al.*, *Proc. Natl. Acad. Sci. USA* 94: 5798-5803 (1997)). Typical LCRs are exemplified by the human β -globin cluster, and the HS-40 regulatory region from the α -globin locus.

25 Eukaryotic vectors may also contain sequences necessary for the termination of transcription and for stabilising the mRNA transcript. Such sequences are commonly available from the 5' and 3' untranslated regions of eukaryotic or viral DNAs, and are known in the art. These regions contain nucleotide segments transcribed as

polyadenylated fragments in the untranslated portion of the mRNA encoding the relevant polypeptide. An appropriate terminator of transcription is fused downstream of the gene encoding the selected nucleic acid binding polypeptide such as a zinc finger protein. Any of a number of known transcriptional terminator, RNA polymerase pause sites and

5 polyadenylation enhancing sequences can be used at the 3' end of the nucleic acid encoding for example a zinc finger polypeptide (see, for example, Richardson, J. P. *Crit. Rev. Biochem. Mol. Biol.* 28:1-30 (1993); Yonaha M. & Proudfoot, N. J. *EMBO J.* 19: 3770-3777 (2000); Ashfield, R. *et al.*, *EMBO J.* 10: 4197-4207 (1991); Hirose, Y. & Manley, J. L. *Nature* 395: 93-96 (1998)).

10 The nucleic acid binding polypeptides are generally targeted to the cell nucleus so that they are able to interact with host cell DNA and bind to the appropriate DNA target in the nucleus and regulate transcription. To effect this, a nuclear localisation sequence (NLS) is incorporated in frame with the expressible nucleic acid binding polypeptide (e.g., zinc finger polypeptide) gene construct. The NLS can be fused either 5' or 3' to the 15 sequence encoding the binding protein, but preferably it is fused to the C-terminus of the chimeric polypeptide.

The NLS of the wild-type Simian Virus 40 Large T-Antigen (Kalderon *et al.* (1984) *Cell* 37: 801-813; and Markland *et al.* (1987) *Mol. Cell. Biol.* 7: 4255-4265) is an appropriate NLS and provides an effective nuclear localisation mechanism in animals. 20 However, several alternative NLSs are known in the art and can be used instead of the SV40 NLS sequence. These include the NLSs of TGA-1A and TGA-1B.

25 Composite binding polypeptides can comprise tag sequences to facilitate studies and/or preparation of such molecules. Tag sequences may include FLAG-tags, myc-tags, 6his-tags, hemagglutinin tags or any other suitable tag known in the art.

Moreover, the nucleic acid binding protein gene according to the invention preferably includes a secretion sequence in order to facilitate secretion of the polypeptide from bacterial hosts, such that it will be produced as a soluble native peptide rather than

in an inclusion body. The peptide may be recovered from the bacterial periplasmic space, or the culture medium, as appropriate.

Construction of vectors employs conventional ligation techniques. Isolated
5 plasmids or DNA fragments are cleaved, tailored, and religated in the form desired to generate the plasmids required. If desired, analysis to confirm correct sequences in the constructed plasmids is performed in a known fashion. Suitable methods for constructing expression vectors, preparing in vitro transcripts, introducing DNA into host cells, and performing analyses for assessing nucleic acid binding protein expression and function
10 are known to those skilled in the art. Gene presence, amplification and / or expression may be measured in a sample directly, for example, by conventional Southern blotting, Northern blotting to quantify the transcription of mRNA, dot blotting (DNA or RNA analysis), or in situ hybridisation, using an appropriately labelled probe which may be based on a sequence provided herein. Those skilled in the art will readily envisage how
15 these methods may be modified, if desired.

f. Applications of Composite Binding Polypeptides

20 Nucleic acid binding proteins according to the invention can be employed in a wide variety of applications, including diagnostics and as research tools, and also in therapeutic applications and in transgenic organisms.

In Vitro Applications

25 Poly-zinc finger peptides of this invention may be employed as diagnostic tools for identifying the presence of nucleic acid molecules in a complex mixture. Nucleic acid binding molecules according to the invention can differentiate single base pair changes in target nucleic acid molecules.

Accordingly, the invention provides methods for determining the presence of a target nucleic acid molecule, wherein the target nucleic acid molecule comprises a target sequence, comprising the steps of:

- 5 a) preparing a nucleic acid binding protein, by a method set forth above, which is specific for the target nucleic acid sequence;
- b) exposing a test system to the nucleic acid binding protein under conditions which promote binding of the protein to the target sequence, and removing any nucleic acid binding protein which remains unbound;
- 10 c) testing for the presence of the nucleic acid binding protein in the test system; wherein, if the nucleic acid binding protein is detected, the target nucleic acid molecule is present and, if the nucleic acid binding protein is not detected, the target nucleic acid molecule is not present. In additional embodiments, quantitation of the amount of nucleic acid binding protein allows quantitation of the amount of the target nucleic acid molecule 15 present in the test system.

In a preferred embodiment, the nucleic acid binding molecules of the invention can be incorporated into an ELISA assay. For example, phage displaying composite binding polypeptides can be used to detect the presence of the target nucleic acid, and visualised 20 using enzyme-linked anti-phage antibodies.

Further improvements to the use of phage expressing a composite binding polypeptide for diagnosis can be made, for example, by co-expressing a marker protein fused to the minor coat protein (gVIII) of a filamentous bacteriophage. Since detection with an anti-phage 25 antibody would then be unnecessary, the time and cost of each diagnosis would be further reduced. Depending on the requirements, suitable markers for display might include fluorescent proteins (A. B. Cubitt, *et al.*, (1995) *Trends Biochem Sci.* **20**, 448-455; T. T. Yang, *et al.*, (1996) *Gene* **173**, 19-23), or an enzyme such as alkaline phosphatase (J. McCafferty, R. H. Jackson, D. J. Chiswell, (1991) *Protein Engineering* **4**, 955-961). 30 Labelling different types of diagnostic phage with distinct markers would allow multiplex screening of a single nucleic acid sample. Nevertheless, even in the absence of such refinements, the basic ELISA technique is reliable, fast, simple and particularly

inexpensive. Moreover it requires no specialised apparatus, nor does it employ hazardous reagents such as radioactive isotopes, making it amenable to routine use in the clinic. The major advantage of the protocol is that it obviates the requirement for gel electrophoresis, and so opens the way to automated nucleic acid diagnosis.

5

The invention provides nucleic acid binding proteins that have exquisite specificity. The invention lends itself, therefore, to the design of any molecule of which specific nucleic acid binding is required. For example, the proteins according to the invention may be employed in the manufacture of chimeric restriction enzymes, in which a nucleic acid cleaving domain is fused to a nucleic acid binding domain comprising a zinc finger as described herein.

10

In Vivo Applications

15

The invention further provides composite binding polypeptides (and nucleic acids encoding them) that may be used in transgenic organisms (such as non-human animals), as therapeutic agents, and in gene therapy applications.

20

A transgenic animal is an animal, preferably a non-human animal, containing at least one foreign gene, called a transgene, in its genetic material. Preferably, the transgene is contained in the animal's germ line such that it can be transmitted to the animal's offspring. Transgenic animals may carry the transgene in all their cells or may be genetically mosaic.

25

Constructs useful for creating transgenic animals according to the invention comprise genes encoding nucleic acid binding polypeptides, optionally under the control of nucleic acid sequences directing their expression in cells of a particular lineage. Alternatively, nucleic acid binding polypeptide encoding constructs may be under the control of non-lineage-specific promoters, and/or inducibly regulated. Typically, DNA fragments on the order of 10 kilobases or less are used to construct a transgenic animal (Reeves, 1998, *New. Anat.*, 253:19). A transgenic animal expressing one transgene can be crossed to a

second transgenic animal expressing second transgene such that their offspring will carry both transgenes.

Although the majority of previous studies have involved transgenic mice, other species of transgenic animal have also been produced, such as rabbits, sheep, pigs (Hammer et al.,

5 1985, *Nature* 315:680-683; Kumar, et al., U.S. 05922854; Seebach, et al., U.S. Patent No. 6,030,833) and chickens (Salter et al., 1987, *Virology* 157:236-240). Transgenic animals are currently being developed to serve as bioreactors for the production of useful pharmaceutical compounds (Van Brunt, 1988, *Bio/Technology* 6:1149-1154; Wilmut, et al., 1988, *New Scientist* (July 7 issue) pp. 56-59). Up-regulation of endogenous or 10 exogenous genes expressing useful polypeptides, such as therapeutic polypeptides, by means of a heterologous nucleic acid binding polypeptide, may be used to produce such polypeptides in transgenic animals. Preferably, the polypeptides are secreted into an extractable fluid, such as blood or mammary fluid (milk), to enable easy isolation of the polypeptide.

15

Furthermore, the invention provides the use of polypeptide fusions comprising an integrase, such as a viral integrase, and a nucleic acid binding protein according to the invention to target nucleic acid sequences *in vivo* (Bushman, (1994) *PNAS* (USA) 91:9233-9237). In gene therapy applications, the method may be applied to the delivery 20 of functional genes into defective genes, or the delivery of a heterologous nucleic acid in order to disrupt an endogenous gene. Alternatively, genes may be delivered to known, repetitive stretches of nucleic acid, such as centromeres, together with an activating sequence such as an LCR. This would represent a route to the safe and predictable incorporation of nucleic acid into the genome.

25

In conventional therapeutic applications, nucleic acid binding proteins according to this embodiment may be used to specifically eliminate cells having mutant vital proteins. For example, if a mutant ras gene is targeted, cells comprising this mutant gene will be destroyed because ras is essential to cellular survival. Alternatively, the action of 30 transcription factors can be modulated, preferably reduced, by administering to the cell

agents which bind to the binding site specific for the transcription factor. For example, the activity of HIV tat may be reduced by binding proteins specific for HIV TAR.

Moreover, binding proteins according to the invention can be coupled to toxic molecules, such as nucleases, which are capable of causing irreversible nucleic acid damage and cell death. Such agents are capable of selectively destroying cells that comprise a mutation in their endogenous nucleic acid.

Nucleic acid binding proteins and derivatives thereof as set forth above may also be applied to the treatment of infections and the like in the form of organism-specific antibiotic or antiviral drugs. In such applications, the binding proteins can be coupled to a nuclease or other nuclear toxin and targeted specifically to the nucleic acids of microorganisms.

Transgenic animals comprising transgenes, optionally integrated within the genome, and expressing heterologous zinc finger and other nucleic acid binding polypeptides from transgenes, may be created by a variety of methods. Methods for producing transgenic animals are known in the art, and are described by Gordon, J. & Ruddle, F.H. *Science* 214: 1244-1246 (1981); Jaenisch, R. *Proc. Natl. Acad. Sci. USA* 73: 1260-1264 (1976); Gossler *et al.*, (1986) *Proc. Natl. Acad. Sci. USA* 83:9065-9069; Hogan *et al.*, *Manipulating the Mouse Embryo: A Laboratory Manual*, (1988); and US. Pat. Nos. 5,175,384; 5,434,340 and 5,591,669.

Pharmaceutical Preparations

The invention likewise relates to pharmaceutical preparations which contain the compounds according to the invention or pharmaceutically acceptable salts thereof as active ingredients, and to processes for their preparation.

The pharmaceutical preparations according to the invention which contain the compound according to the invention or pharmaceutically acceptable salts thereof are those for enteral, such as oral, furthermore rectal, and parenteral administration to (a) warm-

blooded animal(s), the pharmacological active ingredient being present on its own or together with a pharmaceutically acceptable carrier. The daily dose of the active ingredient depends on the age and the individual condition and also on the manner of administration.

5

The novel pharmaceutical preparations contain, for example, from about 10 % to about 80% (or any integral percentage therebetween), preferably from about 20 % to about 60 %, of the active ingredient. Pharmaceutical preparations according to the invention for enteral or parenteral administration are, for example, those in unit dose forms, such as 10 sugar-coated tablets, tablets, capsules or suppositories, and furthermore ampoules. These are prepared in a manner known per se, for example by means of conventional mixing, granulating, sugar-coating, dissolving or lyophilising processes. Thus, pharmaceutical preparations for oral use can be obtained by combining the active ingredient with solid carriers, if desired granulating a mixture obtained, and processing the mixture or granules, 15 if desired or necessary, after addition of suitable excipients to give tablets or sugar-coated tablet cores.

Suitable carriers are, in particular, fillers, such as sugars, for example lactose, sucrose, mannitol or sorbitol, cellulose preparations and/or calcium phosphates, for example 20 tricalcium phosphate or calcium hydrogen phosphate, furthermore binders, such as starch paste, using, for example, corn, wheat, rice or potato starch, gelatin, tragacanth, methylcellulose and/or polyvinylpyrrolidone, if desired, disintegrants, such as the abovementioned starches, furthermore carboxymethyl starch, crosslinked polyvinylpyrrolidone, agar, alginic acid or a salt thereof, such as sodium alginate; 25 auxiliaries are primarily glidants, flow-regulators and lubricants, for example silicic acid, talc, stearic acid or salts thereof, such as magnesium or calcium stearate, and/or polyethylene glycol. Sugar-coated tablet cores are provided with suitable coatings which, if desired, are resistant to gastric juice, using, inter alia, concentrated sugar solutions which, if desired, contain gum arabic, talc, polyvinylpyrrolidone, polyethylene glycol 30 and/or titanium dioxide, coating solutions in suitable organic solvents or solvent mixtures or, for the preparation of gastric juice-resistant coatings, solutions of suitable cellulose preparations, such as acetylcellulose phthalate or hydroxypropylmethylcellulose

phthalate. Colorants or pigments, for example to identify or to indicate different doses of active ingredient, may be added to the tablets or sugar-coated tablet coatings.

Other orally utilisable pharmaceutical preparations are hard gelatin capsules, and also soft 5 closed capsules made of gelatin and a plasticiser, such as glycerol or sorbitol. The hard gelatin capsules may contain the active ingredient in the form of granules, for example in a mixture with fillers, such as lactose, binders, such as starches, and/or lubricants, such as talc or magnesium stearate, and, if desired, stabilisers. In soft capsules, the active 10 ingredient is preferably dissolved or suspended in suitable liquids, such as fatty oils, paraffin oil or liquid polyethylene glycols, it also being possible to add stabilisers.

Suitable rectally utilisable pharmaceutical preparations are, for example, suppositories, which consist of a combination of the active ingredient with a suppository base. Suitable suppository bases are, for example, natural or synthetic triglycerides, paraffin 15 hydrocarbons, polyethylene glycols or higher alkanols. Furthermore, gelatin rectal capsules which contain a combination of the active ingredient with a base substance may also be used. Suitable base substances are, for example, liquid triglycerides, polyethylene glycols or paraffin hydrocarbons.

20 Suitable preparations for parenteral administration are primarily aqueous solutions of an active ingredient in water-soluble form, for example a water-soluble salt, and furthermore suspensions of the active ingredient, such as appropriate oily injection suspensions, using suitable lipophilic solvents or vehicles, such as fatty oils, for example sesame oil, or synthetic fatty acid esters, for example ethyl oleate or triglycerides, or aqueous injection 25 suspensions which contain viscosity-increasing substances, for example sodium carboxymethylcellulose, sorbitol and/or dextran, and, if necessary, also stabilisers.

30 The dose of the active ingredient depends on the warm-blooded animal species, the age and the individual condition and on the manner of administration. For example, an approximate daily dose of about 10 mg to about 250 mg is to be estimated in the case of oral administration for a patient weighing approximately 75 kg .

g. **Transformation and Transfection**

DNA can be stably incorporated into cells or can be transiently expressed using methods known in the art and described below. Stably transfected cells can be prepared by transfecting cells with an expression vector containing a selectable marker gene, and 5 growing the transfected cells under conditions selective for cells expressing the marker gene. To prepare transient transfectants, cells are transfected with a reporter gene to monitor transfection efficiency.

There are many well-known methods of introducing foreign nucleic acids into host cells, which include electroporation, calcium phosphate co-precipitation, particle 10 bombardment, microinjection, naked DNA, liposomes, lipofection, and viral infection etc (see, e.g. Sambrook *et al.* (1989) *Molecular Cloning: A Laboratory Manual*, Second Edition, Cold Spring Harbor Laboratory Press, and Mountain, A. *Trends Biotechnol.* 18: 119-128 (2000) for a review). Any of the above methods can be used, as long as it is compatible with the host cell. Linear nucleic acid molecules have been found to be more 15 efficiently incorporated into mammalian genomes than circular plasmids. Additionally, nucleic acid molecules may be delivered to specific target tissues or to individual cells. Viral based gene transfer is often favoured for introducing nucleic acids into mammalian cells and specific target tissues, and several viral delivery approaches are in clinical trials for gene therapy applications. However, non-viral methods are attractive due to their 20 greater safety for the purpose of gene transfer to humans.

The preferred methods of particle bombardment use biolistics made from gold (or tungsten). Compared with other transfection procedures, particle bombardment requires a low amount of nucleic acid and a smaller number of cells, making the procedure generally more efficient (Heiser, W. C. *Anal. Biochem.* 217: 185-196 (1994); Klein, T. M. 25 & Fitzpatrick-McElligott, S. *Curr. Opin. Biotechnol.* 4: 583-590 (1993)). The procedure is particularly suited for organisms that are difficult to transfect, and for introducing DNA into organelles, such as mitochondria and chloroplasts. Although generally used for *ex vivo* applications, the procedure is also suitable for *in vivo* transfection of skin tissue. Suitable methods are known in the art and described, for instance, in US Patent Nos.

5,489,520 and 5,550,318. See also, Potrykus (1990) *Bio/Technol.* 8: 535-542; and Finnegan *et al.* (1994) *Bio/Technol.* 12: 883-888.

Microinjection is a common method of nucleic acid delivery to isolated cells (Palmiter, R. D. & Brinster, R. L. *Annu. Rev. Genet.* 20: 465-499 (1986); Wall, R. J. *et al.*, *J. Cell Biochem.* 49: 113-120 (1992); Chan, A. W. *et al.*, *Proc. Natl. Acad. Sci. USA* 95: 14028-14033 (1998)). DNA is generally injected into cells and the cells may then be re-introduced into animals. Procedures for such a technique are described in US Pat. Nos. 5,175,384 and 5,434,340, and improvements to the technique are described in WO 00/69257.

10 Efficient for gene transfer *in vivo* can be obtained following local injection of naked DNA. While expression of injected DNA in skin lasts for only a few days, injected DNA in mouse skeletal muscle has been shown to last for up to nine months (Wolff, J. A. *et al.*, *Hum. Mol. Genet.* 1: 363-369 (1992)). Naked DNA is particularly suited to gene therapy for preventive and therapeutic vaccines.

15 Cationic liposomes containing cholesterol are particularly suited for delivery of nucleic acids to humans as they are biodegradable and stable in the bloodstream. Liposomes can be injected intravenously, subcutaneously or inhaled as an aerosol. Stribling *et al.* (1992) *Proc. Natl. Acad. Sci. USA* 89:11,277-11,281. Liposomes can be targeted to certain cell types by incorporating ligands, receptors or antibodies 20 (immunolipids) into the lipid membrane (US. Pat. No. 4,957,773). On contacting target cells, entry of DNA from liposomes is via endocytosis and diffusion. Preparations of lipid formulations are commercially available and methods for their use are well documented (Bogdanenko, E. V. *et al.*, *Vopr. Med. Khim.* 46: 226-245 (2000); Natsume, A. *et al.*, *Gene Ther.* 6: 1626-1633 (1999)).

25 Uptake of DNA into animal cells can also be enhanced by using transfection agents. "Transfected agent", as utilised herein, means a composition of matter added to the genetic material for enhancing the uptake of exogenous DNA segment (s) into a eukaryotic cell, preferably a mammalian cell, and more preferably a mammalian germ

cell. The enhancement is measured relative to the uptake in the absence of the transfecting agent. Examples of transfecting agents include adenovirus-transferrin-polylysine-DNA complexes. These complexes generally augment the uptake of DNA into the cell and reduce its breakdown during its passage through the cytoplasm to the nucleus

5 of the cell. These complexes can be targeted to the male germ cells using specific ligands which are recognised by receptors on the cell surface of the germ cell, such as the c-kit ligand or modifications thereof. Other preferred transfecting agents include lipofectinTM, lipofectamineTM, DIMRIE C, Superfect, and Effectin (Qiagen), unifectin, maxifectin, DOTMA, DOGS (Transfectam; dioctadecylamidoglycylspermine), DOPE (1,2-dioleoyl-

10 sn-glycero-3 phosphoethanolamine), DOTAP (1,2-dioleoyl-3-trimethylammonium propane), DDAB (dimethyl dioctadecylammonium bromide), DHDEAB (N, N-di-n-hexadecyl-N, N-dihydroxyethyl ammonium bromide), HDEAB (N-n-hexadecylN, N dihydroxyethylammonium bromide), polybrene, or poly (ethylenimine) (PEI). For example, Banerjee, R. *et al.*, Novel series of non-glycerol-based cationic transfection

15 lipids for use in liposomal gene delivery, *J. Med. Chem.* 42 (21): 4292-99 (1999); Godbey, W. T. *et al.*, Improved packing of poly (ethylenimine)-DNA complexes increases transfection efficiency, *Gene Ther.* 6 (8): 1380-88 (1999); Kichler, A *et al.*,

Influence of the DNA complexation medium on the transfection efficiency of lipospermine/DNA particles, *Gene Ther.* 5 (6): 855-60 (1998); Birchaa, J. C. *et al.*,

20 Physico-chemical characterisation and transfection efficiency of lipid-based gene delivery complexes, *Int. J. Pharm.* 183 (2): 195-207 (1999). These non-viral agents have the advantage that they facilitate stable integration of xenogeneic DNA sequences into the vertebrate genome, without size restrictions commonly associated with virus-derived transfecting agents.

25 The most critical issues for applications such as gene therapy are the efficient delivery and appropriate expression of transgenes in host cells. For this purpose, viral systems are particularly well suited as viruses have evolved to efficiently cross the plasma membrane of eukaryotic cells and express their nucleic acids in host cells. Suitability of viral vectors is assessed primarily on their ability to carry foreign nucleic acids and 30 deliver and express transgenes with high efficiency. Current applications utilise both RNA and DNA virus based systems, and 70% of gene therapy trials use viral vectors

derived from retroviruses, adenovirus, adeno-associated virus, herpesvirus and pox virus.

See, for example, Flotte *et al.* (1995) *Gene Ther.* 2:357-362; Glorioso *et al.* (1995) *Ann. Rev. Microbiol.* 49:675-710; Smith (1995) *Ann. Rev. Microbiol.* 49:807-838; Prince (1998) *Pathology* 30:335-347; and Robbins *et al.* (1998) *Pharmacol. Ther.* 80:35-47.

5 Retroviruses represent the most prominent gene delivery system as they mediate high gene transfer and expression of therapeutic genes. Members of the DNA virus family such as adenovirus, adeno-associated virus or herpesvirus are popular due to their efficiency of gene delivery. Adenoviral vectors are particularly suited when transient transfection of nucleic acid is preferred. Retroviruses express particular envelope 10 proteins that bind to specific cell surface receptors on host cells, in order for the virus to enter the cell. Hence, the type of viral vector used should be determined by the tissue type to be targeted. See *e.g.*, Dornburg (1995) *Gene Ther.* 2:301-310; Gunzburg, *et al.* (1996) *J. Mol. Med.* 74:171-182; Vile *et al.* (1996) *Mol. Biotechnol.* 5:139-158; Miller (1997) "Development and Applications of Retroviral Vectors" Cold Spring Harbor Laboratory 15 Press, Cold Spring Harbor, New York; Karavanas *et al.* (1998) *Crit. Rev. Oncol. Hematol.* 28:7-30; Hu *et al.* (2000) *Pharmacol. Rev.* 52: 493-511; and Walther *et al.* (2000) *Drugs* 60: 249-271 for reviews.

20 Safety is a critical issue for viral based gene delivery because most viruses are either pathogens or have pathogenic potential. Generally, when a replication-competent virus infects an animal cell it can express viral genes and release many new infectious viral particles in the host organism. Hence, it is very important that during transgene delivery the host animal does not receive a pathogenic virus with full replication potential. For this reason, viral-host cell systems have been developed for gene therapy treatments to prevent the creation of replication-competent viruses. In this method, viral 25 components are divided between a vector and a helper construct to limit the ability of the virus to replicate (Miller 1997). The viral vector contains the gene(s) of interest and cis-acting elements that allow gene expression and replication, but contain deletions of some or all of the viral proteins. Helper cells (or occasionally, helper virus) are engineered to express the viral proteins needed to propagate the viral vectors. These new viral particles 30 are able to infect target cells, reverse transcribe the vector RNA and integrate its DNA copy into the genome of the host, which can then be expressed. However, the vector can

not express the viral proteins required to create new infectious particles. Helper cell lines are known in the art (see Hu, W-S & Pathak, V. K. *Pharmacol. Rev.* 52: 493-511 (2000), for a review).

In general, retroviral vectors are able to package reasonably long stretches of foreign DNA (up to 10 kb). Oncoviruses are a type of retrovirus, which only infect rapidly dividing cells. For this reason they are especially attractive for cancer therapy. Murine leukaemia virus (MLV)-based vectors are the most commonly used of this class. Spleen necrosis virus (SNV), Rous sarcoma virus and avian leukosis virus are other types. Lentiviral vectors are retroviral vectors that can be propagated to produce high viral titres and are able to infect non-dividing cells. They are more complex than oncoviruses and require regulation of their replication cycle. Lentiviral vectors which may be used include human immunodeficiency virus (HIV-1 and -2) and simian immunodeficiency virus (SIV) based systems. HIV infects cells of the immune system, most importantly CD4⁺ T-lymphocytes, and so may be useful for targeted gene therapy of this cell type. Another type of retrovirus is the spumavirus. Spumaviruses are attractive because of their apparent lack of toxicity. Linial (1999) *J. Virol.* 73:1747-1755.

Adenoviral vectors have high transduction efficiency and are able to transfect a number of different cell types, including non-dividing cells. They have a high capacity for foreign DNA and can carry up to 30 kb of non-viral DNA (for a review see, Kochanek, S. *Hum. Gene Ther.* 10: 2451-2459 (1999)). Recombinant adenoviral (rAd) vectors are becoming one of the most powerful gene delivery systems available and have been used to deliver DNA to post-mitotic neurons of the central nervous system (CNS) (Geddes, B. J. *et al.*, *Front. Neuroendocrinol.* 20: 296-316 (1999), and are used to treat diseases such as colon cancer (Alvarez *et al.*, *Hum. Gene Ther.* 5: 597-613 (1997). Adeno-associated virus (AAV) vectors and recombinant AAV (rAAV) vectors are proving themselves to be safe and efficacious for the long-term expression of proteins to correct genetic disease. Snyder, R. O. J. (*Gene. Med.* 1: 166-175 (1999)) provides a review of gene delivery approaches using such vectors. Construction of such vectors is described in, for example, Samulski *et al.*, *J. Virol.* 63: 3822-3828 (1989), and US. Pat. No. 5,173,414.

Many gene therapy trials have been conducted and are underway (over 3,500 people have been treated with gene therapy systems), and several reviews can be studied for details of the protocols and results (Hwu & Rosenberg, *Ann N Y Acad Sci.* 1994 May 31;716:188-97; Blaese, *Hosp Pract (Off Ed)*. 1995 Nov 15;30(11):33-40; Blaese, *Hosp Pract (Off Ed)*. 1995 Dec 15;30(12):37-45; Breau & Clayman, *Curr Opin Oncol.* 1996 May; 8(3):227-31; Dunbar *Annu Rev Med.* 1996;47:11-206; Lotze *Cancer J Sci Am.* 1996 Mar;2(2):63). The first gene therapy trial was carried out by Blaese *et al.*, (1995), to correct a genetic disorder known as adenosine deaminase (ADA) deficiency, which leads to severe immunodeficiency. Several cancer gene therapy strategies are being developed, which involve eliminating cancer cells by suicide therapy (Oldfield *et al.*, *Hum Gene Ther.* 1993 Feb;4(1):39-69), modification of cancer cells to promote immune responses (Lotze *et al.*, *Hum Gene Ther.* 1994 Jan;5(1):41-55), and reversion by delivery of a tumor suppressor gene (Roth *et al.*, *Hum Gene Ther.* 1996 May 1;7(7):861-74). Another successful gene therapy trial has been conducted to combat graft-versus-host disease, which can result following transplant procedures such as bone marrow transplants (Bonini *et al.*, *Science.* 1997 Jun 13;276(5319):1719-24). This procedure was carried out using an HSV-based vector. Several gene therapy treatments are under investigation for the treatment of HIV-1 infection. Most treatments involve modification of lymphocytes, *ex vivo*, to suppress the expression of viral genes, by means of ribozymes, antisense RNA, mutant trans-dominant regulatory proteins and modification to elicit a host immune response (Nabel *et al.*, *Cardiovasc Res.* 1994 Apr;28(4):445-55; Galpin *et al.*, *Hum Gene Ther.* 1994 Aug;5(8):997-1017; Morgan RA, Walker R. *Hum Gene Ther.* 1996 Jun 20;7(10):1281-306 Gene therapy for AIDS using retroviral mediated gene transfer to deliver HIV-1 antisense TAR and transdominant Rev protein genes to syngeneic lymphocytes in HIV-1 infected identical twins; Wong-Staal *et al.*, *Hum Gene Ther.* 1998 Nov 1;9(16):2407-25). Vectors currently in use for gene therapy treatments and animal tests include those derived from Moloney murine leukemia virus, such as MFG and derivative thereof, and the MSCV retroviral expression system (Clontech, Palo Alto, California). Many other vectors are also commercially available.

30 Viral vectors are especially important in applications when a specific tissue type is to be targeted, such as for gene therapy applications. There are two available methods for

targeting genes to specific cell or tissue type. One strategy is designed to control expression of the required gene using a tissue specific promoter (discussed above), and another strategy is to control viral entry into cells. Viruses tend to enter specific cell types according to the envelope proteins that they express. However, by engineering the

5 envelope proteins to express specific proteins as fusions, such as erythropoietin, insulin-like growth factor I and single chain variable fragment antibodies, viral vectors can be targeted to specific cell-types (Kasahara *et al.*, *Science*. 1994 Nov 25;266(5189):1373-6; Somia *et al.*, *Proc Natl Acad Sci U S A*. 1995 Aug 1;92(16):7570-4; Jiang *et al.*, *J Virol*. 1998 Dec;72(12):10148-56; Chadwick *et al.*, *J Mol Biol*. 1999 Jan 15;285(2):485-94).

10 In one example of tissue specific targeting in transgenic mice, a novel transgene delivery system has been developed in which the target tissue type expresses an avian viral receptor (TVA), under the control of a tissue specific promoter. Transgenic mice expressing the TVA receptor are then infected with avian leukosis virus, carrying the transgene(s) of interest (Fisher, G. H. *et al.*, *Oncogene* 18: 5253-5260 (1999)).

15 **h. Construction of Zinc Finger libraries**

Zinc finger libraries may be constructed from naturally-occurring human zinc finger modules. Thus, the invention provides libraries of zinc finger modules. Module libraries according to the invention may be assembled combinatorially into zinc finger polypeptides. The combinatorial assembly may be carried out biologically, using random 20 assembly and selection technologies, or in a directed manner under computer control, assembling desired modules to produce zinc fingers having defined or random specificity. In accordance with the invention, libraries may be constructed entirely from natural zinc finger polypeptide modules from which zinc finger polypeptides having any desired specificity may be isolated. The invention, in its most preferred aspect, does not require 25 the engineering of the specificity of any zinc finger module in order to produce a zinc finger polypeptide having specificity for any desired nucleic acid sequence.

Selection of appropriate zinc finger modules for assembly into libraries of composite binding polypeptides having a predetermined binding specificity can be

accomplished by applying the rules for zinc finger binding specificity set forth herein. In the case of zinc finger assembly under computer control, a rule table may be used to select zinc fingers for binding to the target site. Figure 1 shows a flowchart depicting part of the logic used in the selection of zinc fingers from a natural library in accordance with 5 the invention. The logic set forth in Figure 1 may be supplemented, for example using Rules relating to zinc finger overlap. Functional testing of zinc fingers for binding to the desired binding site may be implemented in an automated fashion and integrated with the zinc finger design system.

10 The invention thus provides libraries of zinc finger modules. In one embodiment, the modules are human zinc finger modules. Preferably, the modules are DNA-binding zinc finger modules.

15 In a preferred aspect the invention provides a library of DNA-binding human zinc finger modules as set out in Example 1 below. Moreover, the invention provides a library of human zinc finger modules as set forth in Example 2 below. Sub-libraries can be prepared from either of the libraries of the invention.

The invention furthermore encompasses libraries in which zinc finger modules as set forth in Examples 1 or 2 herein are combined with other zinc finger modules to provide further libraries that may be used to generate zinc finger polypeptides.

20 In a still further aspect, the invention relates to libraries derived from animals other than humans, for use in said organisms in order to derive some or all of the same advantages as may be obtained with human zinc fingers for use in humans. Example 3 sets forth databases of zinc fingers from mouse, chicken and plants. Sequences of zinc fingers can be identified in other organisms by the same means, *i.e.* by analysis of sequence information and identification of zinc fingers in accordance with the guidance 25 given herein.

EXAMPLES**Example 1. List of selected human DNA-binding zinc fingers.**

These fingers have been selected from the human genome on the basis of a prediction that

5 they have a DNA-binding potential. This prediction is based on coded contacts (WO 96/06166, WO 98/53057, WO 98/53058; WO 98/53059 and WO 98/53060); accordingly, for each peptide unit, a 3-nucleotide DNA target subsite is shown, as the preferred sequence to which the zinc finger binds. Hence, by constructing 2- or 3-finger libraries from these 200 or so units, in the manner described in the Examples *infra*, there 10 exists the potential to screen a large variety of novel DNA target sites. Note that the predicted DNA target subsites listed below are merely intended to be a guide to the DNA-binding potential. It is anticipated that, in practice, an even wider range of DNA sequences can be targeted using a library engineered from this database, through the exertion of a positive selection pressure in the library screening system.

15

The fingers listed below are in a format that can be linked with classical wild-type canonical "TGEKP" (SEQ ID NO:3) linkers (i.e. ...TGEKP - zinc finger peptide sequence - TGEKP - zinc finger peptide sequence - TGEKP - etc...). For each peptide sequence, an oligonucleotide is designed to encode the peptide sequence; the 20 oligonucleotide can then be linked into a library selection system, as described in the Examples *infra*.

Database of predicted human DNA-binding zinc fingers

25 227 finger units

Zinc finger	DNA site	SEQ ID NO	Peptide sequence
ZIF268 F1	GCG	31	YACPVESCDRRFSRSDELTRHIRIH
ZIF268 F2	TGG	32	FQCRICMRNFSRSDELTHIRTH
ZIF268 F3	GCG	33	FACDICGRKFARSDERKRHTKIH
Kr-like13	NGT	34	HKCHYAGCEKVYGKSSHKAHLRTH
MAZ F1	AGG	35	YQCPVCQQRFKRKDRMSYHVRSH

MAZ F2	TGG	36	YNCSHCGKSFSRPDHLSHVRQVH
MAZ F3	NGT	37	FKCEKCEAAFATKDRLRAHTVRH
TIEG2 (SP1) F3	GGG	38	FVCPVCDRRFMRSDDHLSKHARRH
SP1 F1	GGG	39	HKCHYAGCEKVYGKSSHLSKAHLRTH
SP1 F2	GCG	40	FACSWQDCNKKFARSDELARHYRTH
SP1 F3	GGG	41	FSCPICEKRFMRSDDHLSKHARRH
WT1 F1	TGT	42	FMCAYPGCNKRYFKLQLQMHSRKH
WT1 F2	GAG	43	YQCDFKDCERRFSRSRSDQLKRHQRRH
WT1 F3	TGG	44	FQCKTCQRKFSRSRSDHLSKTHTRTH
WT1 F4	GCG	45	FSCRWPSCQKKFARSDELVRHHNMH
TYY1	TAT	46	FQCTFEGCGKRFSLDFNLRTVRIH
TYY1	NAA	47	YVCPFDGCNKFAQSTNLKSHILTH
TF3A	GGG	48	FVCDYEGCGKAFIRDYHLSRSHILTH
TF3A	GGC	49	FKCTQEGCGKHFASPSKLKRHAKAH
MAZ	GGC	50	HACEMCGKAFRDVYHLSRHKLSH
GLI1	GCA	51	YMCEHEGCSKAFSNASDRAKHQNRTH
ZIC3	GCA	52	FKCEFEGCDRRFANSDDRKKHMVH
SP4	NGG	53	HICHIEGCGKVYGKTSHLRAHLRWH
SP2	NTG	54	HVCHIPDCGKTFRKTSLLRAHVRLH
BTE1	NGG	55	HKCPYSGCGKVYGKSSHLSKAHYRVH
GLI2	TAG	56	HKCTFEGCSKAYSRLENLKTHLRSH
Q14872	TAT	57	YQCTFEGCPRTYSTAGNLRTHQKTH
Q14872	TGC	58	FRCDHDGCGKAFAASHHLKTHVRTH
ZIC3	TAG	59	FPCPFPGCGKIFARSENLKIHKRTH
Z143	CTT	60	FKCPFEGCGRSFTTSNIRKVHVRTH
Z143	CGT	61	FRCEYDGCGKLYTTAAHHLKVHERSH
O00153	AAT	62	FMCHESGCGKQFTTAGNLKNHRRIH
Z143	AAC	63	YYCTEPGCGRAFASATNYKNHVRIH
Q14872	TCT	64	FVCNQEGCGKAFLTSHSLRIHVVRH
O00153	TGT	65	FICPAEGCGKSFYVLQRLKVHMRTH
Q14872	GCT	66	FNCESEGCSKYFTTLSDLRKHIRTH
Z143	GCT	67	YRCSEDNCTSKFKTSGDLQKHIRTH
BTE1	GCG	68	FPCTWPDCLKKFSRSDELTRHYRTH
O15391	TAA	69	FVCPFDVCNRKFAQSTNLKTHILTH
Z143	GNC	70	YVCTVPGCDKRFTEYSSLYKHHVVH
O43591	GGT	71	HVCEHCNAAFRTNYHLQRHVFIH
BCL6	TAG	72	YRCNICGAQFNRPANLKTHTRIH
O75626	TAC	73	HECQVCHKRFSSTSNLKTHLRLH
O75626	YAA	74	YECNVCAKTFGQLSNLKVLRLRVH
BCL6	NGA	75	YKCETCGARFVQVAHLRAHVLIH

O75626	GGA	76	FKCQTCNKGFTQLAHLQKHVLVH
ZN45	N (N/T) A	77	YRCDVCGKFRFRQRSYLAHQRVH
BCL6	YTY	78	YPCEICGTRFRHLQTLKSHLRIH
GFI1	GCA	79	YPCQYCGKRFHQKSDMKKHTFIH
Z263	GAN	80	YQCNICGKCFSCNSNLHRHQRTH
ZN75	TAY	81	YRCWSGKSFSHNTNLHTHQRIH
Z186	TTT (YYY)	82	YKCIECGKTFTVNLQHHRTH
Z136	TTT (YYY)	83	FKCKQCGKAFSCSPTLRIHERTH
Z136	TGA	84	YKCKVCGKAFDYPSPRFRTHERSH
Z136	TTT (YYY)	85	YKCKVCGKPFHSLSSFQVHERIH
Z177	TTA	86	YECKECGKAFRNSSCLRVHVRTH
Z136	TNN	87	FECKRCGKAFRSSLSSFRLHERTH
O60765	A/T-YT	88	YRCNECGKGFTSISRLNRHRIIH
ZN42	TYT	89	YHCGECGLGFTQVSRLTEHQRIH
ZN42	CGG	90	FVCGDCGQGFVRSARLEEHRRVH
O14913	TCG	91	YKCEKCGKGFFRSSLQHHQKIH
O14913	C-G/T-G	92	YKCEECGKGFSRSSKLQEHQTIH
ZN45	YYC	93	YKCEECGKGFCRASNLLDHQRGH
ZN45	AAA	94	YKCEECGKGFSQASNLLAHQRGH
ZN45	NAG	95	YQCEECGKGFCRASNFLAHRGVH
Z239	YYG	96	YKCEQCGKGFTRSSSLIHQAVH
O94892	YNY	97	YRCSECGKGFIIVNSGLMLHQRTH
ZN45	AAY	98	YQCAECGKGFSVGSQQLQAHQRCH
ZN45	NGY	99	YKCEECGKGFSVGSHLQAHQISH
ZN45	YCG	100	YQCDACGKGFSRSSDFNIHFRVH
ZN45	CCG	101	YKCGTCGKGFSRSSDLNVHCRIH
ZN45	TGA	102	YKCNACGKSF SYSSHLDNIHCRIH
Z239	TCA	103	YQCYECGKGFSQSSDLRIHLRVH
Z239	YAA	104	YKCGECGKGFSQSSNLHIHRCIH
Z239	YGA	105	YKCDKCGKGFSQSSKLHIIHQRVH
Z239	CGA	106	YHCGKCGKGFSQSSNLHIHQRVH
O60765	AYA	107	FKCSECGRAFSQSASLIQHERIH
O60792	GYY	108	YECKECGKAFIRSSSLAKHERIH
ZN07	ATA	109	YPCKECGKAFSQSSTLAQHQRMH
O43296	AYY	110	YKCSECGKAFSRSSSLTQHQRMH
Z134	ATG	111	YKCSECGKAFSRKDTLVQHQRIH
Z134	ATG	112	YECSECGKAFSRKATLVQHQRIH
ZN84	AYC	113	YECSECGKAFSEKLSLTNHQRIH
Z191	AYG	114	YGCVECGKAFSRSSILVQHQRVH
ZN24	ACG	115	YGCVECGKAFSRSSILVQHQRVH

O43338	GTA	116	YVCQCGKSFQSRATLIKHHRVH
O43339	GTA	117	YECSQCGKSFQKATLKVHQRVH
O43338	AYA	118	YDCQCGKSFQKSSLIQHQVH
O43339	ANA	119	YECQCGKSFQKSGLIQHQVH
O43338	CAA	120	YECGECGKSFQSSNLIEHRIH
Q13398	AAA	121	YECGECGKSFQRSNLMQHRRVH
Z135	CYA	122	YECGECGKAFSQSTLLTEHRIH
Q13398	ACA	123	YECSECGKSFQSSLIQHRRVH
O14709	AAA	124	YKCNECGKAFSQSAYLLNHQRIH
O14709	CAA	125	YKCNECGKVFSQNSAYLIDHQRLH
O14709	CAA	126	YKCTECGKAFQTSAYLFDHQRLH
O14709	CAA	127	YKCDECQKTFQAQTTYLIDHQRLH
O60792	AAA	128	YNCNECRKTFSQSTYLIQHQRIH
O15535	ANA	129	YHCKECGKVFSQSAGLIQHQRIH
Q15776 (a)	TNA	130	YHCKECGKAFSQNTGLILHQRIH
Q15776 (b)	TNA	131	YQCNQCGKAFSQSAGLILHQRIH
Q15776	CNA	132	YKCNECGRAFSQKSGLIEHQRIH
ZN84	AAC	133	YGCNECGRAFSEKSNLINHQRIH
Z191	ANA	134	YKCLECGKAFSQNSGLINHQRIH
ZN24	ANA	135	YKCLECGKAFSQNSGLINHQRIH
O60765	AYA	136	YRCEECGISFGQSSALIQHRRIH
ZN07	YYA	137	YRCEECGKAFGQSSSLIHHQRIH
O43340	ACA	138	YECDECQKSYSQSSALLQHRRVH
Z135	CYY	139	YKCQECGKAFSHSSALIEHHRTH
O43340	AYA	140	YDCSECGKSFQVSVLIQHQRVH
O43340	AYA	141	YVCSECGKSFQKSVLIQHQRVH
Q13398	AYT	142	YQCSQCGKSFQCKSVLIQHQRVH
O15535	GNA	143	HKCDECQKSFQSSGLIRHQRIH
Q15776	GNA	144	HKCDECQKSFQSSGLVRHWRIH
O75802	ANG	145	HKCEECGKAFSRSSGLIQHQRIH
Z189	ANG	146	HKCEECGKAFSRSSGLIQHQRIH
O75802	ANG	147	HKCDECQKAFSRNSGLIQHQRIH
Q13398	YYG	148	HECNECGKSFQSSSLIHHRRLH
Z195	YAA	149	YKCDECQKNTQSSNLIVHKRIH
O43309	CYA	150	YKCDKCGKAFQRSVLTEHQRIH
Z195	CGA	151	YKCDECQKAYTQSSHLEHRIH
ZN45	YYA	152	YKCERCGKAFSQFSSLQVHQRVH
O60893	YYN	153	YECEDCGKTFIGSSALVIHQRVH
ZN07	TAT	154	YECLQCGKAFSMSTQLTIHQRVH
O60893	CYA	155	YECDDCGKTFSQSCSLLEHHKIH

Q15776	NGG	156	YECDECGKTFRRSSHLIGHQRSH
ZN84	YGG	157	YECGECGKAFSRKSHLISHWRTH
Z177	YGA	158	YECDHCGKSFSQLSSHLNVKRTH
O43296	AYG	159	YECMECGKAFNRKSYLTQHQRH
O43296	GNG	160	YECVECGKAFTRMSGLTRHKRIH
O43340	AGG	161	YECRECGKSFTRKNHЛИQHKTВH
Z134	AAG	162	YECSECGKTFSRKDNLTQHKRH
O43338	CGA	163	YECSECGKSFSQTSHLNDHRRH
O75467	AGA	164	YECAQCGKAFSQTSHLTQHQRH
Z135	AGA	165	YECSECGKAFRQSIHLTQHLR
Z135	AGA	166	YECHDCGKSFRQSTHLTQHRRH
Z205	AGG	167	YACTDCGKRFGRSSHЛИQHQIIH
O43296	AGG	168	YECTECGKTFIKSTHLLQHHMIH
O75290	AAG	169	YECKECGKYFSRSANLIQHQSIH
O75290	AGG	170	YECKECGKGFGNREGAHLIQHQKIH
O75290	AGG	171	YECKECGKGFGNREGAHLIQHQKIH
O60792	CGA	172	YTCNECGKAFSQRGHFMEHQKIH
O75123	CGA	173	YTCDQCGKGFGQSSHLMEHQRH
O43337	GYA	174	YECNACGKAQSSTLIRHYLH
O75802	GYY	175	YECNYCGKTFVSSTLIRHQRIH
Z165	GGY	176	YECSECGKTFRVSSHLLIRHFRIH
Z124	CYY	177	YVCNNCGKGFRCSSSLRDHERTH
Z135	AYY	178	YGCNECGKTFSHSSSQHERTH
O15361	GAY	179	YDCNHCGKSFNHKTNLNKHERH
O75123	AAA	180	YVCNECGKRFQTSNFTQHQRIH
Q13398	AYY	181	YVCGEKGKSFSHSSNLKNHQRVH
ZN35	YYA	182	YTCNECGKAQRSSLTVHQRT
Z157	YYC	183	YECTECGKTFSEKATLTIHQRT
O43338	GYY	184	YECDECGKAFTGSKSTLVRHQRT
ZN84	TYC	185	YECSECGKAFTGEKSSLATHHQRT
ZN07	GAA	186	YGCNECGKAFTSQSQLVRHQRT
ZN84	YAA	187	YNCSCQCGKAFTSQSQLTSHQRT
Z186	YGY	188	YACDHCEKAFTSHKSKLTVHQRT
O43338	GGC	189	YVCGEKGKAFTMFKSKLVRHQRT
OZF	YYA	190	YECNVCGKAFTSQSSSLTVHVRSH
O95779	YYY	191	YKCKECGKAFTNHCSLLTIHERTH
Z135	GYY	192	YACRDCGKAFTHSSSLTKHQRT
ZN80	GYA	193	YECKECGKGFGYYSYSLTRHTRSH
Z177	GYC	194	YECSDCGKAFTDQSSLKKHTRSH
Z177	GYY	195	YDCKECGKAFTVPSLQKHVRTH

O43337	ACT	196	YDCMACGKAFRCSSELIOHQRIH
Q14585	AGY	197	YECKECEKAFRSGSKLIQHQRMH
Q14585	AAY	198	YECIDCGKAFGSGSNLTQHRRIH
Q14585	GYY	199	YECKACGMAFSSGSALTRHQRIH
Q14585	AYY	200	YECKECGKAFYSGSSLTQHQRIH
Q14585	AAY	201	YECKECGKAFGSGANLAYHQRIH
Q14585	GAY	202	FECKECGKAFGSGSNLTHHQRIH
Q14585	ACY	203	YVCKECGKAFNSGSDLTQHQRIH
O60792	ACY	204	YQCHECGKTFSYGSSLIQHRKIH
O60893	GNA	205	HYCHECGKSFAQSSGLTKHRRIH
Z165	GCC	206	YECNECGKSFAESSDLTRHRRIH
O60893	GAY	207	YECEECGKVFSHSSNLIKHQRTH
Q15776	NGY	208	YECNECGKAFSHSSHLLIGHHQRIH
Z135	GYY	209	YQCGEKGAFSHSSSLTKHQRIH
Z165	GGY	210	HQCNECGKAFRHSSKLARHQRIH
Z135	TYG	211	YECHECLKGFRNSSALTQHQRIH
O43361	YGC	212	YECNECGKFFLDSYKLVIHQRIH
O43361	YGC	213	YECSECGKFFRDSYKLIIHQRVH
Z140	YYG	214	YGCHECGKTFGRRFSLVLHQRTH
O60792	AAA	215	YECNECGKAFSQHSNLTQHQKTH
Z135	ANA	216	YKCTQCGRTFNQIAPLIHQRTH
Z135	ANA	217	YECNQCGRAFSQLAPLIHQRIH
Z135	ANA	218	YECHECGKAFTQITPLIHQRTH
O43309	AGA	219	YKCNECGKAFGRWSALNQHQRLH
ZN83	AGA	220	YKCNECGKVFNMSHLAQHRRIH
ZN83	AGY	221	YRCNVCGKVFHISHLHQRIH
ZN83	AGA	222	YKCNECGKVFNQISHLHQRIH
O14709	CAY	223	FECSECGRAFSSNRNLIEHKRIH
ZN74	GYA	224	YKCSECGRAFSQNHCLIKHQKIH
Q13398	ANA	225	YECSECGKSFSQNFSLIYHQRVH
O75123	GYA	226	FECKECGKGFSQSSLLIRHQRIH
Z132 (a)	GGA	227	FECSECGRDFSQSSHLLRHQKVH
Z132	GYA	228	YECNECGKFFSQNSILIKHQKVH
Z132 (b)	GGA	229	YECDECGKAFSNRSHLIRHEKVH
Z132	GGN	230	YECSECGRAFSSNSHLVRHQRVH
Z132	AAA	231	YECSECGRAFNNNSNLAQHQKVH
Z134	ATY	232	YKCSDCGKVFRHKSTLVQHESIH
O75290	AAT	233	YECKECGKAFRLYLQLSQHQKTH
Z157	AYC	234	YECGECKNFRAKKSLNQHQRIH
Z157	TTT	235	YECGECKFFRMKMTLNNHQRTH

ZN07	AAT	236	YECAECGVFRQLCSQLNQHQRIH
Z157	AYT	237	YECSECGKIFSMKKSLCQHRRTH
O43361	GGY	238	YECNKCGKFFMYNSKLIRHQKVH
O43361	GTY	239	YKCSKCGKFFRYRCTLRSRHQKVH
Z157	CGY	240	YECNECGNAFYVKARLIEHQRMH
Z157	CGY	241	YECSECGNAFYVKVRLIEHQRIH
O75123	AGG	242	FECNECGKAFIRSSKLIQHQRIH
ZN07	AGT	243	FKCTECGKAFRLSSKLIQHQRIH
O75123	GYT	244	YECNECGKAFFLSSYLLIRHQKIH
O75802	AAT	245	HKCGECGKAFRLSTYLIQHQKIH
Z174	GCG RNA	246	YKCDDCGKSFTWNSELKRHKRVH
Z202	GCG RNA	247	YRCDDCGKHFRWTSIDLVRHQRTTH
O43345	GTG RNA	248	YKCEECGKAYKWPSTLSYHKKIH
O43345	CA? RNA	249	YKCEECGKAFNWSSNLMEHKKIH
O75346	TAA	250	YRCEECGKAFNQSANLTTHKRIH
ZN43	TAA	251	YKCEECGKAFTQSSNLTTHKKIH
ZN85	GGA	252	YKCEECGKAFNQSSKLTKHKKIH
ZN85	GAA	253	YTCEECGKAFNQSSNLTKHKRIH
Q02313	GAA	254	YKCEECGKAFNQLSNLTRHKVIH
Q02313	CAA	255	YKCEECGKAFKQFSNLTDHKKIH
Z141	GTG	256	YKCEECGKAFNRSTTLTKHKRIH
ZN91	TTG	257	YKCEECGKAFSRSSTLTKHKTIH

Example 2: List of all human C₂H₂ zinc fingers

This list represents an even more comprehensive database of human zinc fingers, including those with non-DNA-binding activities such as those mediating protein-protein interactions and those involved in RNA binding. By including fingers from this database

5 into a natural finger selection system as disclosed herein, many new zinc finger proteins having unique target specificities can be obtained. All of these peptides would necessarily possess properties required for potential therapeutic agents, such as non-immunogenicity.

10 The fingers listed below are in a format that can be linked with classical canonical “TGEKP” linkers (i.e. . . . TGEKP – zinc finger peptide sequence – TGEKP – zinc finger peptide sequence – TGEKP - etc...). For each peptide sequence, an oligonucleotide is designed to encode the peptide sequence; the oligonucleotide can then be linked into a library selection system, as described in the Examples *infra*.

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Human zinc finger database**968 finger units**

Name	SEQ ID NO	Peptide sequence
Q92981_HUMAN	258	HQCAHCEKTFNRKDHLKNHFQTH
O76019_HUMAN	259	HQCAHCEKTFNRKDHLKNHLQTH
ZFY_HUMAN	260	HRCEYCKKGFRRPSEKNQHIMRH
ZFX_HUMAN	261	HRCEYCKKGFRRPSEKNQHIMRH
ZFX_BOVIN	262	HRCEYCKKGFRRPSEKNQHIMRH
Q15558_HUMAN	263	HRCEYCKKGFRRPSEKNQHIMRH
ZFX_HUMAN	264	HKCDMCDKGFFHRPSELKKHVAAH
ZFY_HUMAN	265	HKCEMCEKGFFHRPSELKKHVAVH
Q15558_HUMAN	266	HKCEMCEKGFFHRPSELKKHVAVH
Z161_HUMAN	267	YTCSVCVGKGSFSPDHL SCHVKHVH
MAZ_HUMAN	268	YNC SHCGKSF SRFPDHL NSHVRQVH
O43829_HUMAN	269	YSCEVCGKSFIRAPDLKKHERVH
O00403_HUMAN	270	YSCEVCGKSFIRAPDLKKHERVH
Z151_HUMAN	271	HKCPHCDKKFNQVGNLKAHLKIH
Q92618_HUMAN	272	YKCPYCDHRASQKGNLKIHIRSH
ZFX_HUMAN	273	FRCKRCKGFRQQSELKKHMKTH
Q14526_HUMAN	274	YPCTICGKKFTQRGTMTRHMRSH
HKR3_HUMAN	275	FECTECGYKFTRQAHLRRHMEIH
Q14526_HUMAN	276	YACDACGMRFTRQYRLTEHMRIH
O75626_HUMAN	277	YECNVCAKTFGQLSNLK VHLRVH
CTCF_HUMAN	278	HKCPDCDMAFVTSGELVRHRRYKH
O75701_HUMAN	279	YSCPDCSLRFAYTSLLAIHRRIH

075701_HUMAN	280	YACSDCKSRFTYPYLLAIHQRKH
043167_HUMAN	281	YACKDCGKVFKYNHFLAIHQRSRSH
075850_HUMAN	282	CACPDCGRSFTQRAHMLLHQRSRSH
075850_HUMAN	283	YACPDCGRGFSHGQHLARHPRVH
ZN42_HUMAN	284	FVCGDCGQGFVRSARLEEHRRVH
075467_HUMAN	285	FRCVDCGKAFAKGAVLLSHRRIH
015015_HUMAN	286	YKCSECGRAYRHRGSLVNHRHSH
075701_HUMAN	287	YPCPDCGRRFRQRGSLAIHRRAH
Q92951_HUMAN	288	YECAICQRSFRNQSNLAVHRRVH
BCL6_HUMAN	289	YKCDRCQASFRYKGNLASHKTVH
ZN42_HUMAN	290	YACQDCGRRFHQSTKLIQHQRVH
075701_HUMAN	291	YPCPDCGRRFTYSSLLLHSHRRIH
075701_HUMAN	292	HVCTDCGRRFTYPSLLVSHRRMH
075701_HUMAN	293	HSCPDCGRNFSYPSLLASHQRVH
ZN42_HUMAN	294	YACVECGERFGRRSVLLQHRRVH
043298_HUMAN	295	YGCGVCGKKFKMKHHLVGHMKIH
015209_HUMAN	296	YDCPVCNKKFKMKHHLTEHMKTH
043829_HUMAN	297	YACHMCDKAFKHKSHLKDHERRH
000403_HUMAN	298	YACHMCDKAFKHKSHLKDHERRH
060315_HUMAN	299	HQCQICKKAFKHKHHHLIEHSRLH
Q12924_HUMAN	300	HECGICKKAFKHKHHHLIEHMRLH
NIL2_HUMAN	301	HECGICKKAFKHKHHHLIEHMRLH
Q12924_HUMAN	302	FKCTECGKAFKYKHHLKEHLRIH
060315_HUMAN	303	FKCTECGKAFKYKHHLKEHLRIH
NIL2_HUMAN	304	FKCTECGKAFKYKHHLKEHLRIH
095780_HUMAN	305	YKCEECGKAFKRCSHLNEHKRVQ
095779_HUMAN	306	YKCEECGKAFKRCSHLNEHKRVQ
043296_HUMAN	307	FKCSECGKVFNKKHLLAGHEKIH
014709_HUMAN	308	YKCKECGKGFYRHSGLIHLRRH
014709_HUMAN	309	HKCKECGKGFIQRSSLLMHLRNH
ZN80_HUMAN	310	CKCVECGKVFNRRSHLLCYRQIH
043337_HUMAN	311	YKCIECGKAFKRRSHLLQHQRVH
060765_HUMAN	312	YICKECGKAFTLSTSILYKHLRTH
Z136_HUMAN	313	FECKRCGKAFRSSSSFRLHERTH
Z136_HUMAN	314	FVCKQCGKAFRSASTFQIHERTH
Z136_HUMAN	315	YVCKHCGKAFVSSTSIRIHERTH
Z136_HUMAN	316	FKCKQCGKAFSCSPTLRIHERTH
Z124_HUMAN	317	YVCNNCGKGFRCSSSLRDHERTH
Z177_HUMAN	318	YECKECGKAFRNSSCLRHVVRTH
Z124_HUMAN	319	YECKHCGKAFRYSNCLHYHERTH
095780_HUMAN	320	YKCKECGKAFNHCSLLTIHERTH
095779_HUMAN	321	YKCKECGKAFNHCSLLTIHERTH
Z124_HUMAN	322	YPCKQCGKAFRYASSLQKHEKTH
Z136_HUMAN	323	YECKQCGKAFSYLNSFRTHEMIH
Z136_HUMAN	324	YECKQCGKAFSYLPSLRLHERIH
015060_HUMAN	325	YSCKVCGKRFKAHTSEFNYHRRIH
Z136_HUMAN	326	YKCKVCGKPFHSLSPFRIHERTH
Z136_HUMAN	327	YKCKVCGKPFHSLSSFQVHERIH

Z136_HUMAN	328	YKCKVCGKAFDYPSPRFRTHERSH
ZN35_HUMAN	329	YVCNECGKAFTCSSYLLIHQRIH
O15322_HUMAN	330	YNCKECGKSFRWSSYLLIHQRIH
Q92951_HUMAN	331	YRCDQCGKAFSQKGSLIVHIRVH
Q92951_HUMAN	332	YQCKECGKSFSQRGSLAVHERLH
Q92951_HUMAN	333	YECQECGKSFRQKGSLTLHERIH
OZF_HUMAN	334	YECNECGKAFSQRTSLIVHVRIH
OZF_HUMAN	335	YECNVCGKAFSQSSLTVHVRSRSH
ZN07_HUMAN	336	YVCNDCGKAFSQSSSLIYHQRIH
Z151_HUMAN	337	CQCVMCGKAFTQASSLIAHVRQH
Z177_HUMAN	338	YDCKECGKAFTVPSSLQKHVRTH
OZF_HUMAN	339	FECKDCGKAFIQKSNLIRHQRTH
Z177_HUMAN	340	YECSDCGKAFIDQSSLKKHTRSH
Z177_HUMAN	341	YECSDCGKAFIFQSSLKKHMRSH
O60792_HUMAN	342	YECKECGKAFIRSSSLAKHERIH
Z161_HUMAN	343	YACTYCSKAFRDSYHLRRHESCH
Z161_HUMAN	344	HACEMCGKAFRDVYHLMNRHKLSH
MAZ_HUMAN	345	HACEMCGKAFRDVYHLMNRHKLSH
O60792_HUMAN	346	FKCDCEDKTFTRSTHLTQHQKIH
O60792_HUMAN	347	YKCNECDKAFSRSTHLTEHQNTH
Z263_HUMAN	348	YKCNECGKSFRQGMHLTRHQRTH
Z263_HUMAN	349	HKCLECGKCFSQNTHLTRHQRTH
Z135_HUMAN	350	YECSQCGKAFRQSTHLTQHQRIH
Z135_HUMAN	351	YECHDCGKSFRQSTHLTQHRRIH
Z135_HUMAN	352	YECSCECGKAFRQSIHLTQHLRIH
O75467_HUMAN	353	YECAQCGKAFSQTSHTQHQRIH
ZN07_HUMAN	354	YECLQCGKAFSMSTQLTIHQRVH
O95270_HUMAN	355	YPCQFCGKRFHQKSDMKKHTYIH
GFI1_HUMAN	356	YPCQYCGKRFHQKSDMKKHTFIH
O75850_HUMAN	357	FPCTECEKFRKKTHLIRHQRIH
Q15552_HUMAN	358	FRCDECGMRSIQKYHMERHKRTH
O43591_HUMAN	359	FRCDECGMRFIQKYHMERHKRTH
Q15552_HUMAN	360	FQCSQCDMRFIQKYLLQRHEKIH
O43591_HUMAN	361	FQCSQCDMRFIQKYLLQRHEKIH
O75850_HUMAN	362	FPCSECDKRFSSKKAHLTRLRTH
O75850_HUMAN	363	YPCAECGKRFSSQKIHLSHQKTH
O94892_HUMAN	364	FMCSECGKGFTMKRYLIVHQQIH
O43336_HUMAN	365	YQCSECGKSFIYKQSLLDHHRIH
O43167_HUMAN	366	FKCNECGKGFAQKHSLOVHTRMH
O43167_HUMAN	367	YTCDQCGKYFSQNRQLKSHYRVH
PLZF_HUMAN	368	YECNGCDKKFSLKHQLETHYRVH
HKR3_HUMAN	369	YACPTCHKKFLSKYYLKVNKRKH
O43336_HUMAN	370	YVCNVCGKSFRHKQTFVGHQQRIH
O43336_HUMAN	371	YVCNICGKSFLHKQTLVGHQQRIH
Z134_HUMAN	372	YDCSDCGKSFGHKYTLIKHQRIH
Z200_HUMAN	373	YDCNHCGKSFNHKTNLNKHERIH
O15361_HUMAN	374	YDCNHCGKSFNHKTNLNKHERIH
ZN84_HUMAN	375	YDCNHCGKAFSRKSQLVRHQRTH

ZN84_HUMAN	376	FECRECGKAFSRKSQLVTHHRTH
ZN07_HUMAN	377	YGCRECGKAFSQSQLVRHQRTH
ZN84_HUMAN	378	YRCIECGKAFSQSQLINHQRTH
ZN84_HUMAN	379	YGCSECRKAFSQSQLVNHQRIH
ZN84_HUMAN	380	HGCIQCGKAFSQKSHLISHQMTH
ZN84_HUMAN	381	YNCSQCGKAFSQSQLTSHQRTH
ZN84_HUMAN	382	YVCSECGKAFSQKSHLISHQRTH
Z157_HUMAN	383	FECNECGKSFGRKSQLLILHRTTH
ZN84_HUMAN	384	FECSECGKAFSRKSHLIPHQRTH
ZN84_HUMAN	385	YECGEKGAFSRKSHLISHWRTH
Z136_HUMAN	386	YHCKECGKAYSCRASFQRHMLTH
Z136_HUMAN	387	YECKECGEAFSCIIPSMRRHMIKH
Z136_HUMAN	388	YECQEKGKAFTCITSVRRHMIKH
ZN80_HUMAN	389	YECQEKGKAFPEKVDVVRHMRIH
O43338_HUMAN	390	YVCGEKGKAFMFKSKLVRHQRTH
O43338_HUMAN	391	YECDECCKAFGSKSTLVRHQRTH
Z133_HUMAN	392	YACGEGRGFSQKSNLVAHQRTH
Z133_HUMAN	393	YMCSECGRGFSQKSNLIIHQRTH
Z133_HUMAN	394	YACKDCGRGFSQQSNLIRHQRTH
Z133_HUMAN	395	YACSDCGLGFSDRSNLISHQRTH
Z133_HUMAN	396	YACRECGRGFNRKSTLIIHERTH
Z133_HUMAN	397	YVCRECGRGFSHQAGLIRHKRKH
Z133_HUMAN	398	CVCRECGQGFLQKSHLTLHQMTTH
Z133_HUMAN	399	YVCRECGKGFSQLSAVVRHQRTH
O94892_HUMAN	400	YICSECCKGFPKSNLIVHQRNH
O94892_HUMAN	401	YICNECGKGFPKRNLIIVHQRNH
O94892_HUMAN	402	YTCSECCKGKGPLKSRLIVHQRTTH
O94892_HUMAN	403	YICSECCKGKFTTKHYVIHQRNH
O94892_HUMAN	404	YICSECCKGFTGKSMLIIHQRTH
O94892_HUMAN	405	YLCSECCKGFTVKMSMLIIHQRTH
O94892_HUMAN	406	YGCNECGKGFTMKSRLIVHQRTH
O94892_HUMAN	407	YICNECGKGFTMKSRMIEHQRTH
O94892_HUMAN	408	FICSECCKVFTMKSRLIEHQRTH
O94892_HUMAN	409	YICNECGKGFAFKSNLVVHQRTH
Z186_HUMAN	410	YECNECGKTFHQKSFLTIVHQRTH
Z186_HUMAN	411	YECNELGKTFHCKSFLTIVHQKTH
Z186_HUMAN	412	YGCNECGKTVRCKSFLTTLHQRTH
ZN35_HUMAN	413	YTCNECGKAFRQRSSLTVHQRTH
Z186_HUMAN	414	YQCSECGKTFSQLSYLTIHHRTH
Z157_HUMAN	415	YECSECGKTFRVKISLTQHHRTTH
Z186_HUMAN	416	YKCIECGKTFVNQLLTLHHRTH
Z157_HUMAN	417	YECTECGKTFSEKATLTIHQRTH
ZN84_HUMAN	418	YACSDCRKAFFEKSSELIRHQTIH
ZN84_HUMAN	419	YECSLCRKAFFEKSSELIRHLRTTH
Z140_HUMAN	420	YECNECRKALRCHSFLIKHQRIH
ZN84_HUMAN	421	YECNECRKAFREKSSLINHQRIH
ZN84_HUMAN	422	YECSECRKAFRERSSLINHQRTH
ZN84_HUMAN	423	YECSECGKAFGEKSSLATHQRTH

ZN84_HUMAN	424	YECSECGKAFSEKLSLTNHQRIH
O43339_HUMAN	425	YECSECGKAFRGKYSLVQHQRVH
Z157_HUMAN	426	YECSECGKIFSMKKSLCQHRRTH
Z157_HUMAN	427	YECGECKFFRMKMTLNHHQRTH
Z157_HUMAN	428	YECGECKNFRAKKSLSNQHQRIH
O43361_HUMAN	429	YKCSECGKAFSLKHNVVQHLKIH
Z134_HUMAN	430	YECSECGKAFSRKATLVQHQRIH
Z134_HUMAN	431	YKCSECGKAFSRKDTLVQHQRIH
Z134_HUMAN	432	YECSECGKTFSRKDNLTQHKRIH
O14709_HUMAN	433	YKCCECGKVFIRSKSLLLHQRVH
O14709_HUMAN	434	YECDECGKCFILKKSILGHQRIH
O14709_HUMAN	435	YECNECGKVFILKKSILHQRFH
O14709_HUMAN	436	YKCNKCQKAFILKKSILHQRIH
Z140_HUMAN	437	YACAECDKAFSRSFSLILHQRTH
Z140_HUMAN	438	YGCHECGKTFGRRFSLVLHQRTH
O95878_HUMAN	439	YACAQCGKTFNNTSMLRTHQRIH
O14709_HUMAN	440	YKCDMCCCKHFMKISHLINHRRIH
ZN83_HUMAN	441	FKCDICGKIFNKKSNLASHQRIH
ZN07_HUMAN	442	HQCEDCEKIFRWRSHLIIHQRIH
Z137_HUMAN	443	HKCDDCGKVLTSRSHLIRHQRIH
Z140_HUMAN	444	HECKDCNKTFSYLSFLIEHQRTH
Z189_HUMAN	445	HKCSDCGKAFSWKSHLIEHQRTH
O75802_HUMAN	446	HKCSDCGKAFSWKSHLIEHQRTH
O14709_HUMAN	447	YKCNDGKVFSYRSNLIAHQRIH
O43309_HUMAN	448	YGCDDCGKAFSQHSHLIEHQRIH
O75123_HUMAN	449	YTCDQCGKGFGQSSHLMEHQRIH
O43336_HUMAN	450	YNCTACEKAFIYKNKLVEHQRIH
O43309_HUMAN	451	YKCDVCEKAFIQRTSLTEHQRIH
O60792_HUMAN	452	YKCDQCGKGFIEGPSLTQHQRIH
O43309_HUMAN	453	YKCDKCGKAFTQRSVLTEHQRIH
ZN91_HUMAN	454	YKCEECGKAFKQLSTLTTKRIH
ZN91_HUMAN	455	YKCKECKGKAFQFSTLTTKIIIH
ZN91_HUMAN	456	YKCKECDKTFKRLSTLTKHIIIH
ZN91_HUMAN	457	YKCKECDKTFKRLSTLTKHIIIH
ZN85_HUMAN	458	YKCEKCGKAFNHFSSHLLTTKIIIH
ZN85_HUMAN	459	YKCEECGKAFNRFSTLTTKIIIH
ZN43_HUMAN	460	YKCEECGKAFNQFSTLTKHIIIH
ZN43_HUMAN	461	YTCEECGKVFNWSSRLTTKRIH
ZN43_HUMAN	462	YKCEECGKAFNKSSILTTKIIIR
O75437_HUMAN	463	YKWEKFGKAFNRSSHLLTDKITH
O43345_HUMAN	464	YKCEEEGGKAFNWSSTLTYYKSAH
ZN91_HUMAN	465	YKCEECGKAFNQSSNLTTKIIIH
ZN91_HUMAN	467	YKCEECGKAFNRSSKLTTKIIIH
Q02313_HUMAN	468	YKCEECGKAFNQSSTLTTHNIIIH
ZN91_HUMAN	469	YKCEECGKAFNHSSSLSTHKIIIH
ZN43_HUMAN	470	YKCEECGKAFKLSSTLSTHKIIIH
ZN91_HUMAN	471	YKCEECGKAFQSSTLTTKIIIH
Q02313_HUMAN	472	YKCEECGKAFNQSSTLTTKRIH

O95780_HUMAN	473	YKCEECGKAFNSSSILTEHKVIH
O95779_HUMAN	474	YKCEECGKAFNSSSILTEHKVIH
ZN91_HUMAN	475	YKCKECGKAFKHSSALAKHIIH
ZN85_HUMAN	476	YKCKECGKAFKHSSTLTKHIIH
ZN85_HUMAN	477	YKCEECDKAFKWSSVLTCHKIIH
ZN43_HUMAN	478	YKCEECGKAFKWSSTLTKHIIH
ZN85_HUMAN	479	YKCEECGKGFKWPSTLTIHKIIH
ZN91_HUMAN	480	YKGECGKAFKESSALTCHKIIH
ZN91_HUMAN	481	YKCEECGKAFRKSSTLTEHKIIH
ZN91_HUMAN	482	YKCEECGKAFRQSSTLTKHIIH
Q02313_HUMAN	483	YKGECGKAFNQSSALNTHIIH
ZN91_HUMAN	484	CKCKECEKTFHWSSTLTNHKEIH
O75437_HUMAN	485	YKCKECGKTFNWSSTLTNHRKIY
ZN91_HUMAN	486	YKCKECGKAFSNSSTLANHKITH
ZN91_HUMAN	487	YKCKECGKAFSNSSTLANHKITH
O43345_HUMAN	488	YKCKECGKTFIKVSTLTTHKAIH
O43345_HUMAN	489	YKCEECGKTFSKVSTLTTHKAIH
O43345_HUMAN	490	YKCEECGKTFSKVSTLTTHKAIH
O43345_HUMAN	491	YKCEECGKAFSKVSTLTTHKAIH
O43345_HUMAN	492	YKCKECGKAFSKVSTLITHKAIH
O95270_HUMAN	493	YACRMCGKAFKRSSTLSTHLLIH
GFI1_HUMAN	494	YDCKICGKSFKRSSTLSTHLLIH
O75346_HUMAN	495	YKCIICGKAFKRSSTLTTHKKIH
ZN43_HUMAN	496	YKCKECGKAFNQYSNLTTHNKIH
ZN85_HUMAN	497	YKCKECGKAFNRSSLTTHRKIH
ZN91_HUMAN	498	YKCSEECDKAFIWSSTLTEHKIH
ZN91_HUMAN	499	YKCEECGKAFISSSTLNGHKRIH
ZN43_HUMAN	500	YKCEECGKAFNYSSHNLTHKRIH
O95780_HUMAN	501	YKCEECGKAFNWSSILTEHKRIH
O95779_HUMAN	502	YKCEECGKAFNWSSILTEHKRIH
O43345_HUMAN	503	YKCEECGKAFNWSSNLMEHKRIH
O43345_HUMAN	504	YKCEECGKAFNWSSNLMEHKRIH
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ZN91_HUMAN	507	FKCKECGKAFIWSSTLTRHKRIH
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ZN91_HUMAN	509	YKCEECGKAFLWSSTLRRHKRIH
ZN91_HUMAN	510	YKCEECGKAFLWSSTLTRHKRIH
Q02313_HUMAN	511	YKCEAYGRAFWNSSTLNHKRIH
ZN91_HUMAN	512	YKFEECGKAFRQLTLNKHIIH
Z141_HUMAN	513	YKCEECGKAFRRSTDRSQHKKIH
O75346_HUMAN	514	YKCEECGKAFNWSSDLNKHKKIH
ZN91_HUMAN	515	YKCEECGKAFNWSSSLTKHKRIH
ZN91_HUMAN	516	YKCEECGKAFNWSSSLTKHKRFH
ZN85_HUMAN	517	YKCEECGKAFNWSSLTCHKRIH
ZN43_HUMAN	518	YKCEECGKAFNWPSTLTKHNRIH
ZN43_HUMAN	519	YKCEECGKAFNWPSTLTCHKRIH
O75437_HUMAN	520	YKCEECGKAFFWSSTLTKHKRIH

O95780_HUMAN	521	YKCEECGKAFNWCSSLTKHKRIH
O95779_HUMAN	522	YKCEECGKAFNWCSSLTKHKRIH
ZN43_HUMAN	523	YKCEECGKAFSRSSNLTKHKKIH
ZN43_HUMAN	524	YKCTECGEAFSRSSNLTKHKKIH
ZN91_HUMAN	525	YKCEECGKAFSRSSNLTKHKTIH
O75437_HUMAN	526	YKCEECGKAFNRSSTFTKHVIIH
Z141_HUMAN	527	YKCEECGKAFNRFTTLTKHKRIH
Z141_HUMAN	528	YKCEECGKAFNRSTTLTKHKRIH
ZN43_HUMAN	529	CKCEKCGKAFNCPSIITKHKRIN
O43345_HUMAN	530	YKCEACGKAYNTFSILTKHKVIIH
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O43345_HUMAN	533	YKCEECGKSFSTFSVLTKHKVIIH
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O95780_HUMAN	542	YKCEKCDKVFKRFSYLTKHKRIH
O95779_HUMAN	543	YKCEKCDKVFKRFSYLTKHKRIH
O95780_HUMAN	544	CICEECGKTFKWFSYLTKHKRIH
O95779_HUMAN	545	CICEECGKTFKWFSYLTKHKRIH
ZN43_HUMAN	546	YKCEECGKAFNHFSILTKHKRIH
ZN91_HUMAN	547	YKCEKCKAQNQSSILTNHKKIH
Q02313_HUMAN	548	YKCEKCVRAFNQASKLTEHKLIH
ZN85_HUMAN	549	YKSKECEKAFNQSSKLTEHKKIH
ZN43_HUMAN	550	YKCKECAKAFNQSSNLTEHKKIH
ZN85_HUMAN	551	YKCEECGKAFNQSSKLTKHKKIH
ZN85_HUMAN	552	YKCEECGKAFNQSSNLHKKIH
O43345_HUMAN	553	YKCEECGKAFNRSAILIKHKRIH
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O43345_HUMAN	555	YKCEECGKAFNQSAILTKHKIIH
ZN43_HUMAN	556	YKCEVCGKAFNQFSNLTTHKRIH
ZN43_HUMAN	557	YTCEECPKAFNQFSNLTTHKRIH
O75346_HUMAN	558	YRCEECGKAFNQSANLTTHKRIH
ZN85_HUMAN	559	YTCEECPKAFNQSSNLTKHKRIH
Z141_HUMAN	560	YKCKDCDKAFKRFSHLNHKKIH
Z141_HUMAN	561	YKCKECDKAFKQFSLLSQHKKIH
Q02313_HUMAN	562	YKCEECGKAFKQFSNLTDHKKIH
ZN43_HUMAN	563	YKCEECGKAFTQSSNLTTHKKIH
ZN43_HUMAN	564	YKCEECGKAFTQSSNLTTHKKIH
ZN85_HUMAN	565	YKCEECGKAFQSSNLTTHKIIH
Q02313_HUMAN	566	YKCEECGKAFNQLSNLTRHKVIIH
ZN85_HUMAN	567	YECEKCGKAFNQSSNLTRHKKSH
O95780_HUMAN	568	YNCEECGKAFNRCSHLTRHKKIH

O95779_HUMAN	569	YNCEECGKAFNRCSHLTRHKKIH
O95780_HUMAN	570	YTCEDCGRAFNRSHTKHKTIH
O95779_HUMAN	571	YTCEDCGRAFNRSHTKHKTIH
Q02313_HUMAN	572	YECEEKGKAFNRSSKLTEHKYIH
ZN91_HUMAN	573	YKCEECGKAFNRSSNLTIHKFIH
ZN91_HUMAN	574	YKCEECGKAFNRSSNLTIHKFIH
ZN43_HUMAN	575	YKCEKGKAFNRPSNLIEHKKIH
Z141_HUMAN	576	YTCEECKIIFTSSNFAHKRIH
Z141_HUMAN	577	FTCEECSIFTTSSHFAHKIIH
Z141_HUMAN	578	YTCEEKGKAFKWSLIFNEHKRIH
Z141_HUMAN	579	YTCEEKGKAFRQSSKLNEHKVH
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O43345_HUMAN	586	YKCEECGKAFSWLSVFSKHKKIH
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O95779_HUMAN	589	YKCEECGKAFHWCSPLFVRHKKIH
Z195_HUMAN	590	YTCEECKNIFKQLSDLTKHKKTH
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O43345_HUMAN	592	YKCEECGKAFWPSRLTEHKATH
O43345_HUMAN	593	YKCEECDKAFSWPSSLTEHKATH
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ZN43_HUMAN	595	YKCEECGKAFKWSSKLTEHKLTH
ZN91_HUMAN	596	YKCEECGKAFSHSSALAKHRIH
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ZN74_HUMAN	604	YKCDECGKAFTWSTNLLEHRRIH
Z195_HUMAN	605	YKCDECGKAYTQSSHLEHRRIH
Z195_HUMAN	606	YKCDECGKNFTQSSNLIVHKRIH
Z195_HUMAN	607	YKCDECGKNFTQSSNLIVHKRIH
ZN80_HUMAN	608	YKCKECGSVFNKNSLLVRHQQIH
Z165_HUMAN	609	FGCKECGRAFNLSNLIRHQRIH
Q02313_HUMAN	610	YKCKECGKAFNQTSHLIRHKRIH
O60792_HUMAN	611	YKCNECGRAFNQNIHLTQHKRIH
ZN74_HUMAN	612	YRCGECGKAFNQRTHLTRHRIH
Q15776_HUMAN	613	YKCKECGKAFNGNTGLTQHLRIH
O43309_HUMAN	614	YKCDECGNAFRGITSLIQHQRIH
O43309_HUMAN	615	YKCEECGKAFRGRTVLIRHKIIH
O75123_HUMAN	616	YVCNECGKRFQTSNFTQHQRIH

O60792_HUMAN	617	YKCNECGKAFNGPSTFIRHHMIH
O43296_HUMAN	618	FVCSECGKAFTHCSTFILHKRAH
O43337_HUMAN	619	YECSQCRKAFTHRSTFIRHNRTH
O43296_HUMAN	620	YKCNECGKAFTHRSNFVLHNRRH
OZF_HUMAN	621	YGCNECGKAFSQFSTLALHLRIH
ZN83_HUMAN	622	YKCNERGKAFHQGLHLPIHQIIH
ZN07_HUMAN	623	YKCNECGKAFSQNSTLFQHQIIH
ZN83_HUMAN	624	YKCNECGKVFSRNSYLAQHLIIH
ZN83_HUMAN	625	YECNCGKVFSRNSYLVQHLIIH
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ZN83_HUMAN	629	YKCNECGKVFHNQISHLQAQHQRIH
ZN83_HUMAN	630	YRCNVCGKVFHHISHLQAQHQRIH
ZN83_HUMAN	631	YKCDECGKVFQSQNSYLAHWRIH
Z189_HUMAN	632	YKCDECGKTFVSVAHLVQHQRIH
O75802_HUMAN	633	YKCDECGKTFVSVAHLVQHQRIH
ZN83_HUMAN	634	YKCDECDKAQFSQNSHLVQHHRIH
O60792_HUMAN	635	YKCDECGKAFSQRTHLVQHQRIH
O43361_HUMAN	636	YECESSKVFKYNSSLIKHQIIH
ZN83_HUMAN	637	FKCNECGKAFSMRSSLTNHHAIH
O60792_HUMAN	638	YKCNECGKAFSYCSSLTQHRRIH
Z137_HUMAN	639	YKYHDCGKVFQSASSYAKHRRIH
O14709_HUMAN	640	YKCEDCGKAFSYNSLLVHRRIH
Z124_HUMAN	641	YVCMECGKAFSCLSSLQGHIAH
O60792_HUMAN	642	YQCHECGKTFSYGSSLIQHRKIH
O60792_HUMAN	643	YDCAECGKSFSYWSSLAQHLKIH
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ZN83_HUMAN	645	YKCNECGKVFSHKSSLVNHRIH
Z132_HUMAN	646	YKCSECGKFFSRKSSLICHWRVH
O43339_HUMAN	647	YKCNECGKFFSQTSHLNDHRRIH
O43338_HUMAN	648	YECSECGKSFQTSQSHLNDHRRIH
ZN45_HUMAN	649	YKCNACGKSFSYSSHNIHCRIH
ZN45_HUMAN	650	YKCGTCGKGFSRSSDLNVHCRIH
Z263_HUMAN	651	YKCPLCGKNFSNNNSNLIRHQRIH
Z202_HUMAN	652	YTCPTCGKSFSRGYHLIRHQRTH
O75850_HUMAN	653	FSCPQCGKSFSRKTHLVRHQLIH
Z205_HUMAN	654	YACPLCGKSFSSRRSNLHRHEKIH
O15535_HUMAN	655	HQCIECGKSFNRHCNLRHQKIH
ZN24_HUMAN	656	YECVQCGKSYSQSSNLFRHQRRH
Z191_HUMAN	657	YECVQCGKSYSQSSNLFRHQRRH
Q99592_HUMAN	658	YTCTQCGKSFQYSHNLSRAVVH
Q13397_HUMAN	659	YTCTQCGKSFQYSHNLSRAVVH
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O75802_HUMAN	661	YLCRQCGKSFSQLCNLIRHQGVH
Z189_HUMAN	662	YQCKECGKSFSQLCNLTRHQRIH
O75802_HUMAN	663	YQCKECGKSFSQLCNLTRHQRIH
Z263_HUMAN	664	YKCTLGENFSHRSNLIRHQRIH

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O95878_HUMAN	666	YKCSECGKSFSRSSNRIRHERIH
Z263_HUMAN	667	YTCHECGDSFSHSSNRIRHLRTH
O43336_HUMAN	668	YVCIIICGKSFIRSSDYMQRHQRIH
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BCL6_HUMAN	670	YRCNICGAQFNRPANLKTHTRIH
Z133_HUMAN	671	YKCGECGLSFSKMTNLLSHQRIH
ZN75_HUMAN	672	YRCSWCGKSFSHNTNLHTHQRIH
O60893_HUMAN	673	YKCNECERSFTRNRSLIEHQKIH
ZN74_HUMAN	674	YKCSECGRAFSQNHCLIKHQKIH
O14709_HUMAN	675	YACSECGKGFTYNRNLIEHQRIH
Z177_HUMAN	676	YKCFQCEKAFTSTNLIMHKRIH
O60792_HUMAN	677	YKCNECEKAFSRSENLINHQRIH
O94892_HUMAN	678	YGCTLCAKVFSRKSRLNEHQRIH
Z189_HUMAN	679	YHCTKCKKSFSRNSLLVEHQRIH
O75802_HUMAN	680	YHCTKCKKSFSRNSLLVEHQRIH
O43309_HUMAN	681	YQCTQCNKSFSSRSILTQHQGVH
O15535_HUMAN	682	YQCSQCSKSYSRRSFLIEHQRSH
Z205_HUMAN	683	YTCPACRKSFSHHSTLHQRIH
Z189_HUMAN	684	YTCIECGKSFSSSFLIEHQRIH
O75802_HUMAN	685	YTCIECGKSFSSSFLIEHQRIH
Z189_HUMAN	686	FQCNECGKSFSSSFLIEHQRIH
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O14709_HUMAN	690	YECHVCRKVLTSSRNLMVHQRIH
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ZN35_HUMAN	692	YECNECGKTFTRSSNLIVHQRIH
O75123_HUMAN	693	YECNECGKSFIRSSSLIRHYQIH
O43296_HUMAN	694	YECVECGKSFCKWSTNLIRHAIIH
O43296_HUMAN	695	YECSECGKVFLESAALIHHYVIH
O43337_HUMAN	696	YECTQCGKAFHRSTYLIQHSVIIH
O43296_HUMAN	697	YECTECGKTFIKSTHLLQHHMIH
O75290_HUMAN	698	YECKECGKYFSRSANLIQHQSIH
Z205_HUMAN	699	YACTDCGKRGFRSSHLIQHQIIH
Z165_HUMAN	700	YECSECGKTFRVSSHLIRHFRIH
Q15776_HUMAN	701	YECDECGKTFRRSSHLIGHQRSH
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O75802_HUMAN	704	YECNYCGKTFVSSTLIRHQRIH
O43337_HUMAN	705	YECNACGKAFSQSSTLIRHYLIH
ZN07_HUMAN	706	YECSECGKAFSRSSSYLIEHQRIH
Z132_HUMAN	707	YECSECGKAFAHSSSTLIEHWRVH
O43340_HUMAN	708	YECSECGKAFSCNIYLIHHQRFH
Z135_HUMAN	709	YECGEKGKAFTSQSTLLTEHRRIH
O43338_HUMAN	710	YECGEKGKSFSQSSNLIEHCRIH
O43338_HUMAN	711	YECGKCGKSFTQHSGLILHRKSH
Z140_HUMAN	712	YECDECGKVFTWHASLIQHTKSH

Q13398_HUMAN	713	YACPECGKSFSQIYSLNSHRKVKH
Q13398_HUMAN	714	YECSKCGKSFKQSSSFSSHRKVKH
O43340_HUMAN	715	YECSECCKSFSHSTNLFRHWRVH
O43340_HUMAN	716	YECSECCKSFSHSTNLYRHRSAH
O43340_HUMAN	717	YECSECCKSFSQSSGLLRHRRVH
O43340_HUMAN	718	YKCSECCKSFSQSSGFLRHRKAH
O43340_HUMAN	719	YECSECCKVFSQSSGLFRHRRRAH
O43340_HUMAN	720	YECDECCKSFSQSSALLQHRRVH
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Q13398_HUMAN	722	YECGECGKSFSQRSNLMQHRRVH
Z132_HUMAN	723	YECSECRKSFSRSSSLIQHWRIH
Z132_HUMAN	724	YECSQCGKSFSRSSLALIQHWRVH
Q13398_HUMAN	725	HECNECGKSFSRSSLALIHRRRLH
O43339_HUMAN	726	YKCGECGNSFSQSAILNQHRRIH
O43339_HUMAN	727	YKCGDCGKSFSQSSILIQHRRIH
O60765_HUMAN	728	YRCEECGISFGQSSALIQHRRIH
O43338_HUMAN	729	YECGQCGKSFSLKCGLIQHQLIH
O43339_HUMAN	730	YECGQCGKSFSQKSGLIQHQVVH
O43338_HUMAN	731	YDCGQCGKSFIQKSSLIQHQVVH
Q13398_HUMAN	732	YQCSQCGKSFQCKSVLIQHQRVH
O43340_HUMAN	733	YVCSECCKSFGQKSVLIQHQRVH
O43340_HUMAN	734	YDCSECCKSFRQVSVLIQHQRVH
Q13398_HUMAN	735	YECSECSKSFSCKSNLIKHLRVH
O43339_HUMAN	736	YEKGQCGKSFSQKATLIKHQRVH
O43338_HUMAN	737	YVCQCGKSFSQRATLIKHHRVH
O43339_HUMAN	738	YECSQCGKSFSQKATLVKHQRVH
Q13398_HUMAN	739	YECSECGKSFSQNFSLIYHQRVH
O43340_HUMAN	740	YECSVCGKSFIRKTHLIRHQTVH
O43340_HUMAN	741	YECSECEKSFSCKTDLIRHQTVH
O43340_HUMAN	742	YECRECGKSFRTRKNHLIQHKTVH
Z189_HUMAN	743	HKCEECGKGFGVRKAHFIQHQRVH
O75802_HUMAN	744	HKCEECGKGFGVRKAHFIQHQRVH
O43340_HUMAN	745	HECSECCKSFSRKTHLTQHQRVH
O43309_HUMAN	746	YQCKECGKSFSQSGLIQHQRIH
Q15776_HUMAN	747	YQCNQCGKAFSQSAGLILHQRIH
O15535_HUMAN	748	YHCKECGKVFSQSAGLIQHQRIH
O60792_HUMAN	749	YNCNECRKTFSQSTYLIQHQRIH
Q15776_HUMAN	750	YHCKECGKAFSQNTGLILHQRIH
ZN84_HUMAN	751	YGCNECGRAFSEKSNLINHQRIH
Q15776_HUMAN	752	YKCNECGRAFSQKSGLIEHQRIH
Z189_HUMAN	753	HKCDECGKAFSRNSGLIQHQRIH
O75802_HUMAN	754	HKCDECGKAFSRNSGLIQHQRIH
Z189_HUMAN	755	HKCEECGKAFSRSSGLIQHQRIH
O75802_HUMAN	756	HKCEECGKAFSRSSGLIQHQRIH
ZN24_HUMAN	757	YKCLECGKAFSQNSGLINHQRIH
Z191_HUMAN	758	YKCLECGKAFSQNSGLINHQRIH
OZF_HUMAN	759	YQCSECGKAFSQKSHHIRHQKIH
Q15776_HUMAN	760	YQCNECGKAFIQRSSLIRHQRIH

ZN35_HUMAN	761	YDCSECGKAFSQLSSLIVHQRIH
ZN07_HUMAN	762	YRCEECGKAFGQSSSLIHHQRIH
O60765_HUMAN	763	FKCNTCGKTFRQSSSRIAHQRIH
OZF_HUMAN	764	FKCSECGTAFGQKKYLIKHQNIH
OZF_HUMAN	765	FECNECGKAFSQKQYVIKHQNTH
Q92951_HUMAN	766	FECTHCGKSFRAGKGNLVTHQRIH
OZF_HUMAN	767	FECNECGKSFSDKENLLTHQKIH
ZN74_HUMAN	768	FKCNECGKAFSSHAYLIVHRRIH
ZN74_HUMAN	769	FKCADCGKGFSCAYLLVHRRIH
O60765_HUMAN	770	FKCSECGRAFSQSASLIQHERIH
ZN35_HUMAN	771	FECHECGKAFIQSANLVVHQRIH
ZN35_HUMAN	772	FTCSVCGKGFQSANLVVHQRIH
ZN35_HUMAN	773	FACNDCGKAFTQSANLIVHQRSW
O14709_HUMAN	774	YKCNECGKDFSQNKNLVVHQRMH
O14709_HUMAN	775	YKCDECGKTFAQTTYLIDHQRLH
O14709_HUMAN	776	YKCNECGKVFSQLNAYLIDHQRLH
O14709_HUMAN	777	YKCTECGKAFTQSAYLFDHQRLH
O14709_HUMAN	778	YKCNECGKAFSQSAYLLNHQRIH
Z157_HUMAN	779	YQCNECGKFSRVHSSLGIHQRIH
O60765_HUMAN	780	YNCNECGKALSSHSTLIIHERIH
EVI1_HUMAN	781	YKCDQCPKAFNWKSNLIRHQMSH
Q15776_HUMAN	782	YQCNVCGKAFSYRSALLSHQDIH
O43309_HUMAN	783	YECNECGKAFVYNSSLVSHQEIH
Z200_HUMAN	784	YGCKKGRRFGRLSNCTRHEKTH
O15361_HUMAN	785	YGCKKGRRFGRLSNCTRHEKTH
ZN07_HUMAN	786	YKCNDCGKAFNRSSRLTQHQKIH
ZN74_HUMAN	787	YQCGSCGKAFTCHSSLTVHEKIH
ZN35_HUMAN	788	YVCSKCGKAFTQSSNLTVHQKIH
Z140_HUMAN	789	YECIECGKAFRRFSHLTRHQSIH
O60893_HUMAN	790	YQCNMCGKAFRRNSHLLRHQRIH
Q13396_HUMAN	791	YSCTECEKSFVQKQHLLQHQKIH
O43361_HUMAN	792	YECTQCAKAFVRKSHLVQHEKIH
O43361_HUMAN	793	YECTECEKAFVRKSHLVQHQKIH
O75123_HUMAN	794	YECKECGKAFLQKAHLTEHQKIH
O75290_HUMAN	795	YECKECGKGFNRGGAHLIQHQKIH
O75290_HUMAN	796	YECKECGKGFNRGGAHLIQHQKIH
O75290_HUMAN	797	FECKECGKAFLHMQLIRHQKLH
O75290_HUMAN	798	FECKECGKAFLHMHLIRHQKLH
O75290_HUMAN	799	FECKECGKAFLHIQFTRHQKFH
O75290_HUMAN	800	YECKECGKAFLYLQLSQHQKTH
Z140_HUMAN	801	YECTECGKAFSRASNLTQHQRIH
O43296_HUMAN	802	YECVECGKAFTRMSGLTRHKRIH
O43296_HUMAN	803	YECMECGKAFNRKSYLTQHQRIH
O14913_HUMAN	804	HECVECGKRFSSSRLQEHHQKIH
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O15535_HUMAN	806	YECNECGKAFSRSSGLFNHRGIH
Z132_HUMAN	807	YECNDCGKAFSNSSTLIQHQKVH
Z132_HUMAN	808	YECIQCGKAFSERSTLVRHQKVH

Z132_HUMAN	809	YECDECGKAFSNRSHLIRHEKvh
Z124_HUMAN	810	YECQKCGKAFSRASLWKHKKTH
ZN35_HUMAN	811	FKCNECEKAFSYSSQLARHQKVH
O60792_HUMAN	812	FECSECGKAFSYLSNLNQHQKTH
O75467_HUMAN	813	FRCSECGKAFSHGSNLSQHRKIh
O75467_HUMAN	814	FACPCGCGRAFSHSSNLTOHQOLLH
OZF_HUMAN	815	FACKVCGKVFSHKSNLTEREHFH
Z132_HUMAN	816	YECSQCGKLFSHLCNLAQHKKIh
O60765_HUMAN	817	YECNTCGKLFNHRSSLTNHYKIh
O60792_HUMAN	818	YECAECGKAFRHCSSLAQHQKTH
O43336_HUMAN	819	CECSECGKCFRHRRTSLIQHQKVH
O43336_HUMAN	820	CECNECGKVFSHQKRLLEHQKVH
O95878_HUMAN	821	YECTECGRTFSDISNFGAHQRTH
O60792_HUMAN	822	YECNECGKAFSQHSNLTQHQKTH
O43309_HUMAN	823	YHCNDGKAFSQKAGLFHHIKIH
O43336_HUMAN	824	YECSDCGKAFISKQTLKHHKIh
O60893_HUMAN	825	YECDDCGKTFSQSCSLLHHKIh
O43338_HUMAN	826	FECDECGKSFSQRTTLNKHHKVH
O75123_HUMAN	827	YVCSYCGKGFQRSNFIQHQKIH
O60792_HUMAN	828	YTCNECGKAFSQRGHFMEHQKIH
ZN42_HUMAN	829	YTCDVCGKVFQRSNLLRHQKIH
O14709_HUMAN	830	YGCNDCSKVFRQRKNLTVHQKIH
O43361_HUMAN	831	YVCSECGKAFLTQAHLDGHQKIQ
O43361_HUMAN	832	YTCSECGKAFLTQAHLVGHQKIH
O43361_HUMAN	833	YECTQCGKAFLTQAHLVGHQKTH
Z157_HUMAN	834	YECGECAKTFSARSYLIAHQKTH
O75123_HUMAN	835	YECNECGKAFFLSSYLIHQKIH
Q13398_HUMAN	836	YECNECGKFTYYSSFIHQRVH
O43361_HUMAN	837	YKCSKCGKFFRYRCTLSRHQKVH
O43361_HUMAN	838	YECNCGKFFMYSNKLIRHQKVH
Z132_HUMAN	839	YECNECGKFFSQNSILIKHQKVH
Q13396_HUMAN	840	YECGYCGKSFSHPSDLVRHQRIH
O75467_HUMAN	841	YACPVCGKAFRHSSLVRHQRIH
Z165_HUMAN	842	HQCNECGKAFRHSSKLRHQRIH
Z205_HUMAN	843	YHCLDCGKSFSHSSHLTAHQRTH
Z135_HUMAN	844	YACRDCGKAFTHSSSLTKHQRTH
Z135_HUMAN	845	YECNDGKAFSHSSSLTKHQRIH
Z135_HUMAN	846	YQCGECGKAFSHSSSLTKHQRIH
ZN74_HUMAN	847	FDCSQCWKAFCFSCHSSLIMHQRIH
ZN74_HUMAN	848	YTCGECGKAFSCHSSLNVHQRIH
ZN35_HUMAN	849	YECKECGKAFSCFSHLIVHQRIH
O43309_HUMAN	850	YKCNECGKAFGRWSALNQHQRIH
ZN24_HUMAN	851	YGCVECGKAFSRSSILVQHQRVH
Z191_HUMAN	852	YGCVECGKAFSRSSILVQHQRVH
O43296_HUMAN	853	YKCSECGKAFSRSSSLTOHQRMH
ZN75_HUMAN	854	FKCQECGKSFRVSSDLIKHHRIH
O75290_HUMAN	855	FVCKECGMAFRYHYQLIEHCQIH
O75467_HUMAN	856	FVCTQCGRAFRERPALFHHQRIH

ZN74_HUMAN	857	FKCEKCGEMFNWSSHLTEHQRLH
ZN85_HUMAN	858	FKCTKCGKSGMISCLTEHSRIH
ZN43_HUMAN	859	FKCCECGKSFMLPHLAQHKIIH
Z195_HUMAN	860	FKCQECGKSFQMLSFLTEHQKIH
ZN07_HUMAN	861	FKCDECGKAFRWISRLSQHOLIH
Z189_HUMAN	862	HKCCECGKAFRLSTYLIQHQKIH
O75802_HUMAN	863	HKCCECGKAFRLSTYLIQHQKIH
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O75290_HUMAN	867	FECRECGKAFSLLNQLNRHKNIH
O75290_HUMAN	868	FECKCEKAFSNRAHLIQHYIIH
O43296_HUMAN	869	FECKECGKAFSNRDLIRHFSIH
O62425_CAEEL	870	FVCKVCGKAFRQASTLCRHKIIH
O75123_HUMAN	871	FECKDCGKAFIQSSKLLLHQIIH
O75290_HUMAN	872	FECKECGKFFRQGSNLNQHRSIH
O75290_HUMAN	873	FECKECGKSFNRRSSNLVHQSIH
O75290_HUMAN	874	FECKECGKSFNRRSSNLVHQSIH
O75290_HUMAN	875	FECQDCGKAFNRGSSLVHQSIH
O94892_HUMAN	876	FVCSECRKAFSSKRNLLIVHQRTH
O14709_HUMAN	877	FECSECGRAFSSRNLLIEHKRIH
Z135_HUMAN	878	YECNQCGRASARATLLIEHQRIH
Z157_HUMAN	879	FECQECGKAFCRKAHLTEHQRTH
Z157_HUMAN	880	FECNECGKAYCRKSNLVEHRLIH
O75123_HUMAN	881	FECNECGKAFIRSSKLIQHQRIH
ZN42_HUMAN	882	FRCAECGQSFRQRSNLLHQRIH
ZN42_HUMAN	883	FACPECQGSFRQHANLTQHRRIH
ZN42_HUMAN	884	FACAECGQSFRQRSNLTQHRRIH
ZN42_HUMAN	885	--CAECGKAFRQRPTLTQHLRVH
ZN42_HUMAN	886	YACPECQKAFRQRPTLTQHLRTH
O14913_HUMAN	887	YKCEECGNSFYYPAMLKQHQRIH
Z174_HUMAN	888	YTCGECGNCFGRQSTLKLHQRIH
PLZF_HUMAN	889	YECEFCGSCFRDESTLKSHKRIH
BCL6_HUMAN	890	YPCEICGTRFRHLQTLKSHLRIH
O43296_HUMAN	891	FECLECGKAFNHRSYLKRHQRIH
O43337_HUMAN	892	YKCLECGKAFKRRSYLMQHHPIH
O43296_HUMAN	893	YECLECGKVFKHRSYLMWHQQT
O75123_HUMAN	894	YECKECGKAFRHRSDLIEHQRIH
O43336_HUMAN	895	YECKECGKAFIHKKRLLEHQRIH
Z157_HUMAN	896	YECSECGNAFYVKVRLIEHQRIH
Z157_HUMAN	897	YECNECGNAFYVKARLIEHQRMH
OZF_HUMAN	898	FVCKECGKTFSGKSNLTEHEKIH
Z134_HUMAN	899	YKCSDCGKVFHKSTLVQHESIH
O60893_HUMAN	900	YECEDCGKTFIGSSALVIHQRVH
O43339_HUMAN	901	YECSECGKLFRQNSSLVDHQKIH
O43338_HUMAN	902	FECSECGKFFRQSYTLVEHQKIH
O43338_HUMAN	903	YECGEKGKLFRQSFSLVVHQRIH
O43361_HUMAN	904	YECSECGKLFMDSFTLGRHQRVH

O43361_HUMAN	905	YECSECGKFFRDSYKLIHQRVH
O43361_HUMAN	906	YECNECGKFFLDSYKLVIHQRIH
O43336_HUMAN	907	YECSECGKGFILEVKLLQHQRIH
ZN07_HUMAN	908	YECAECGVFRQLCSQLNQHQRIH
Z132_HUMAN	909	HVCKECGKAFSHSSKLRKHQKFH
TYY1_HUMAN	910	HVCAECGKAFVESSKLKRHQLVH
O15391_HUMAN	911	HVCAECGKAFLESSKLRRHQLVH
O94892_HUMAN	912	HVCSECGKAFVKKSQLTDHERVH
ZFX_HUMAN	913	HICVECGKGFRHPSELKKHMRIH
ZFY_HUMAN	914	HICVECGKGFRYPSELRKHMRIH
Q15558_HUMAN	915	HICVECGKGFRHPSELRKHMRIH
Z135_HUMAN	916	YECHECLKGFRNSSALTQHQRIH
ZN74_HUMAN	917	YTCGECGKAFRQSSSLTLHRRWH
Z174_HUMAN	918	YQCGQCGKSFQSSNLHQHRLH
Z195_HUMAN	919	YQCEECGVVRTCSSLNSHKRTH
HKR3_HUMAN	920	FQCHLCGKTFRTQASLDKHNRTH
O43337_HUMAN	921	YDCMACGKAFRCSSELIQHQRIH
O60765_HUMAN	922	YLCNECGNTFKSSSSLRYHQRIH
O60765_HUMAN	923	YKCNECGKTFRCNSSLSNHQRIH
Z140_HUMAN	924	YKCNECGKAFSSGSELIRHQITH
Q14585_HUMAN	925	YECKECGKAFSGSGLIRHQIIH
Q14585_HUMAN	926	YICNECGKAFSGSALTRHQRIH
Q14585_HUMAN	927	YECKECGKSFGSSGSALNRHQRIH
Q14585_HUMAN	928	YECKACGMAFSSGSALTRHQRIH
Q14585_HUMAN	929	YECKECGKSFSFESALIRHHIH
Q14585_HUMAN	930	YECKECGKTFSGSQSLTQHHRIH
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Q14585_HUMAN	932	YECKECGKAFYSGSSLTQHQRIH
Q14585_HUMAN	933	FECKECGKAFSGSNSLTHHQRIH
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Q14585_HUMAN	936	YECKECGKAFGSGSKLIQHQLIH
Q14585_HUMAN	937	YECKECEKAFRSGSKLIQHQRMH
ZN80_HUMAN	938	YECKECGKTFFYNNSSLTRHMKIH
ZN80_HUMAN	939	YECKECGKGFFYSYSLTRHTRSH
Z165_HUMAN	940	YECNECGKSFAESSDLTRHRRIH
Z202_HUMAN	941	YKCTICGKSFSQKSVLTTHQRIH
O43167_HUMAN	942	YTCEICGKSFTAKSSLQTHIRIH
Q92618_HUMAN	943	HTCCICGKSFPFQSSLSQHMRKH
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O15535_HUMAN	945	HKCDECGKSFTQSSGLIRHQRIH
O60893_HUMAN	946	HYCHECGKSFAQSSGLTKHRRIH
ZN24_HUMAN	947	HICDECGKHFSQLSALILHQRIH
Z191_HUMAN	948	HICDECGKHFSQLSALILHQRIH
Z140_HUMAN	949	YACKECGKTFSQIISNLVKHQMIH
Q14585_HUMAN	950	YECKECGKDFSFVSVLVRHQRIH
O75123_HUMAN	951	FECKECGKGFSQSSLLIRHQRIH
UKLF_HUMAN	952	FKCNHCDRCFSRSDHLALHMKRH

O95600_HUMAN	953	FRCTDCNRSFSRSDHLSLHRRRH
SP2_HUMAN	954	YACACQCKRFMRSDHLSKHVKTH
SP4_HUMAN	955	YACPECSKRFMRSDHLSKHVKTH
O60402_HUMAN	956	YACPECSKRFMRSDHLSKHVKTH
O75411_HUMAN	957	YACPMCDRRFMRSRSDHLSKHARRH
Q13118_HUMAN	958	YACPMCDRRFMRSRSDHLSKHARRH
O14901_HUMAN	959	YACPVCDDRFFMRSRSDHLSKHARRH
BTE1_HUMAN	960	YACPLCEKRFMRSDHLSKHARRH
SP2_HUMAN	961	FVCNWFFCGKRFTRSDELQRHARTH
SP4_HUMAN	962	FICNWMFCGKRFTRSDELQRHRRTH
O60402_HUMAN	963	FICNWMFCGKRFTRSDELQRHRRTH
EZF_HUMAN	964	YHCDWDGCGWKFARSDELTRHYRKH
O95600_HUMAN	965	YKCTWDGCSWKFARSDELTRHFRKH
UKLF_HUMAN	966	YKCSWEGCEWRFARSDELTRHYRKH
EKLF_HUMAN	967	YACTWEGCGWRFARSDELTRHYRKH
BTE2_HUMAN	968	YKCTWEGCDWRFARSDELTRHYRKH
O14901_HUMAN	969	FNCSDWGCDKKFARSDELSRHRRTTH
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BTE1_HUMAN	972	FPCTWPDCCLKFSRSDELTRHYRTH
EGR4_HUMAN	973	FACPVESCVRSFARSDELNRLRIH
EGR2_HUMAN	974	YPCPAEGCDRRFSRSDELTRHIRIH
EGR1_HUMAN	975	YACPVESCDRRFSRSDELTRHIRIH
EGR3_HUMAN	976	HACPAEGCDRRFSRSDELTRHLRIH
Q16256_HUMAN	977	YQCDFKDCERRFFRSRSDQLKRHQRRH
WT1_HUMAN	978	YQCDFKDCERRFSRSRSDQLKRHQRRH
Q15881_HUMAN	979	YQCDFKDCERRFSRSRSDQLKRHQRRH
Q15881_HUMAN	980	FQCKACQRKFSRSRSDHLKTHTRTH
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EGR1_HUMAN	986	FQCRICMRNFSRSRSDHLTHIRTH
EVI1_HUMAN	987	YTCRYCGKIFPRSANLTRHLRTH
O95878_HUMAN	988	YRCTVCGKHFSSRNLLIRHQKTH
Z140_HUMAN	989	YVCKVCNKSFSWSSNLAKHQRTTH
O60893_HUMAN	990	YECEECGVFSHSSNLIKHQRTTH
Z135_HUMAN	991	YECSECGKSFSSRFSQHERTH
O95878_HUMAN	992	YICCECGKSFSSSSFGVHRTTH
ZN80_HUMAN	993	CKCSECGKTFTYRSVFRHSMTH
ZN80_HUMAN	994	YECSECGKTFSYHSVFIQHRVTH
Z135_HUMAN	995	YGCNECGKSFSSHSSLSQHERTH
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Z202_HUMAN	999	CRCNECGKSFSSRDHLVLRHQRTTH
ZN74_HUMAN	1000	FKCSDCEKAFNSRSRLTLHQRTTH

ZN42_HUMAN	1001	FACPECGQRFSQRLKLTRHQERTH
Z205_HUMAN	1002	YPCPECGKCFQSRSNLIAHNRTTH
ZN75_HUMAN	1003	FKCDECCKRFIQNSHLIKHQERTH
ZN07_HUMAN	1004	FKCDECCKGFVQGSHLIQHQRIH
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Z135_HUMAN	1009	YKCQECGKAFSHSSALIEHHRTTH
O60765_HUMAN	1010	FKCKECSKAFSQSSALIQHQITH
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O60792_HUMAN	1012	CKCNECGKAFSYCSALIRHQERTH
Z151_HUMAN	1013	YVCERCGKRFVQSSQLANHIRHH
EVI1_HUMAN	1014	YECENCAKVFTDPSNLQRHIRSQH
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Z205_HUMAN	1016	HKCPICAKCFTQSSALVTHQERTH
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Z200_HUMAN	1018	YTCPLCGKQFNESSYLIISHQORTH
O15361_HUMAN	1019	YTCPLCGKQFNESSYLIISHQORTH
ZN07_HUMAN	1020	YKCNKCTKAFGCSSRLIRHQERTH
Z263_HUMAN	1021	YQCNICGKCFSCNSNLHRHQERTH
Q13134_HUMAN	1022	YKCELCPYSSSQKTHLTRHMRTH
Q13127_HUMAN	1023	YKCELCPYSSSQKTHLTRHMRTH
CTCF_HUMAN	1024	FQCSLCSYASRDTYKLKRHMRTH
Q99592_HUMAN	1025	YTCSLCGKTFSCMYTLKRHERTH
Q13397_HUMAN	1026	YTCSLCGKTFSCMYTLKRHERTH
Q60765_HUMAN	1027	YKCSLCEKTFINSSLRKHEKNH
ZN74_HUMAN	1028	YKCSACEKAFCSSLLSMHLRVH
ZN75_HUMAN	1029	YKCQQCDRRFRWSSDLNKHFMTTH
Z189_HUMAN	1030	YQCNQCKQSFSQRRLSVKHQRIH
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O75066_HUMAN	1036	YACQYCDAVFAQSIELSRHVRTH
O95878_HUMAN	1037	YRCDICGKSFQSATLAVHHRTH
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Z133_HUMAN	1039	YACKECGRCFRQRTTLVNHQERTH
Z133_HUMAN	1040	YVCVGCGHSFSQNSTLISHRRTH
O43336_HUMAN	1041	YVCIECGKSLSSKYSVLSVEHQERTH
O75467_HUMAN	1042	YACAQCGRRFCRNSHLIQHERTH
Z124_HUMAN	1043	YECKQCGKAFSRSSHLDHERTH
Z177_HUMAN	1044	YEQNQCGKSFSTGSYLVHKRTTH
Z177_HUMAN	1045	YECDHCGKSFSQLSHLNVHKRTTH
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Z135_HUMAN	1048	YEQNQCGRAFSQLAPLQLHQRIH

Z135_HUMAN	1049	YKCTQCGRTFNQIAPLIQHQRTH
O60893_HUMAN	1050	YQCDTCGKGFRRTSYLVQHQRSW
O43337_HUMAN	1051	YKCKQCGKGFRNRKWYLVRHQRVH
Z205_HUMAN	1052	YRCEQCGKGFSWHSHLVTHRRTH
Z202_HUMAN	1053	YRCDDCGKHFRWTSSDLVRHQRTW
ZN45_HUMAN	1054	YRCDVCGKFRQRSYLQAHQRVH
ZN45_HUMAN	1055	YQCDACGKGFSRSSDFNIHFRVH
Z239_HUMAN	1056	YQCYECGKGFSQSSDLRIHLRVH
Z239_HUMAN	1057	YKCDKCGKGFSQSSKLHIHQRTW
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Z239_HUMAN	1059	YKCGECGKGFSQSSNLHIHRCIH
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ZN07_HUMAN	1062	YPCKECGKAFSQSSTLAQHQRMH
Z133_HUMAN	1063	YVCKTCGRGFSLKSHLSRHRKTH
Z133_HUMAN	1064	YVCGVCGRGFSLKSHLNRHQNIH
Z133_HUMAN	1065	YVCGVCEKGFSLKSLARHQKAH
EVI1_HUMAN	1066	YRCKYCDRSFSISSLNLQRHVRNIH
RRE1_HUMAN	1067	YKCQTCERTFTLKHSLSVRHQRIH
O75850_HUMAN	1068	YACAQCGRRFSRKSHLGRHQAVH
O75850_HUMAN	1069	HACAVCARFSKTSKTNLVRHQAIH
O75850_HUMAN	1070	YQCAQCARSFTHKQHLSVRHQRTW
ZN42_HUMAN	1071	FVCSECGRFSRSSHLLRHQLTH
Z132_HUMAN	1072	FECSECGRDFSQSSHLLRHQKVH
ZN35_HUMAN	1073	YECEKCGAAFIINSHLMRHHRTW
Z132_HUMAN	1074	YECSECGRAFSSNSHLVRHQRTW
Z202_HUMAN	1075	YKCMECGKSYTRSSHLLARHQKVH
Z134_HUMAN	1076	YECSECGRKAYSLSSHLLNRHQKVH
Z239_HUMAN	1077	YECSKCGKGFSQSSNLHSHHQRTW
Z165_HUMAN	1078	YECSECGRAFSQSSNLSQHQRIH
Z132_HUMAN	1079	YECSECGRAFNNNSNLAQHQKVH
Z239_HUMAN	1080	YECEECEGMSFSQRSNLHIIHQRDH
O00153_HUMAN	1081	HQCQVCGKTFSQSGSRNVHMRKH
Q13398_HUMAN	1082	YVCGECKFSFSHSSNLKNHQRTW
O15322_HUMAN	1083	YKCEICGKSFCLRSSLNRHYMVH
O75123_HUMAN	1084	FKCAQCGKAFCHSSDLIRHQRTW
O14913_HUMAN	1085	YKCEECDKAFLYHSFLRRHKAVH
O14913_HUMAN	1086	YKCEECDKAFLHHSYLRKHQAVH
ZN83_HUMAN	1087	FKCNECGKLFRDNSYLVHQRFH
O15322_HUMAN	1088	HTCNECGKSFCYISALRIHQRTW
O60792_HUMAN	1089	FGCNDCGKSFYRYSALNKHQRLH
Z137_HUMAN	1090	YKCNKCGKIYRHSYLAZYQRTW
O75123_HUMAN	1091	YVCNVCGKDFIYHSGLIEHQRTW
Z134_HUMAN	1092	YKCNKCGKYFSHHSNLIVHQRTW
O43361_HUMAN	1093	FECSICGKFFSHRSTLNMHQRTW
Z134_HUMAN	1094	FECIECGKFFSRSSDYIAHQRTW
Z134_HUMAN	1095	FVCSKCGKDFIRTSHLVRHQRTW
O14913_HUMAN	1096	YKQECGKSFYRSYLRHYRMH

Z174_HUMAN	1097	YKCDDCGKSFTWNSELKRHKRVH
O60765_HUMAN	1098	YRCKECGKSFSRRSGLFIHQKIH
O43167_HUMAN	1099	YSCGICGKSFSDSSAKRRHCILH
O43829_HUMAN	1100	FVCEMCTKGFTTQAHLKEHLKIH
O00403_HUMAN	1101	FVCEMCTKGFTTQAHLKEHLKIH
O75626_HUMAN	1102	FKCQTCNKGFTQLAHLQKHVLVH
O15322_HUMAN	1103	FKCEQCGKGFRCRAILQVHCKLH
BCL6_HUMAN	1104	YKCETCGARFVQVAHLRAHVLIH
Z195_HUMAN	1105	YKCEKCGKAFTQFSHLTVHESIH
ZN85_HUMAN	1106	YKCKKCGKAFNQSAHLTTHEVIH
Z239_HUMAN	1107	YKCEKCGKGFTRSSSLLIHHAVH
Z239_HUMAN	1108	YKCEQCGKGFTRSSSLLIHQAVH
O15322_HUMAN	1109	YKCEECGKGFTDSLDLHKHQI IH
O15322_HUMAN	1110	YICEKCGRAFIHDLKLQKHQI IH
O14913_HUMAN	1111	YKCEKCGKGFFRSSDLQHHQKIH
O14913_HUMAN	1112	YKCEECGKCFSSFTSLKRHQI IH
O14913_HUMAN	1113	YPYKCEECGKGFSRSSKLQEHQTIH
ZN45_HUMAN	1114	YKGEHCVKSFSWSSHLQINQRAH
ZN45_HUMAN	1115	YKCEECGKGFSWSSSLI IHQRVH
ZN45_HUMAN	1116	YKCEECGKVFSSWSSYLAHQRVH
ZN45_HUMAN	1117	YKCEKCDNAFRRFSSLQAHQRVH
ZN45_HUMAN	1118	YKCERCGKAFSQFSSLQVHQRVH
ZN45_HUMAN	1119	YKCEECGVGFSQRSYLVHLKVH
ZN45_HUMAN	1120	YKCEECGKSFSWRSRLQAHERIH
ZN45_HUMAN	1121	YKCEECGKGFSVGSHLQAHQISH
ZN45_HUMAN	1122	YQCAECGKGFSVGSQSQLQAHQRCH
ZN45_HUMAN	1123	YQCEECGKGFCRASNFLAHRGVH
ZN45_HUMAN	1124	YKCEECGKGFSQASNLLAHQRGH
ZN45_HUMAN	1125	YKCEECGKGFSQASNLLAHQRGH
O75467_HUMAN	1126	FVCALCGAAFSQGSSLFKHQRVH
ZN42_HUMAN	1127	YHCGECGLGFTQVSRLTEHQRIH
O60765_HUMAN	1128	YRCNECGKGFTSISRLNRHRI IH
TYY1_HUMAN	1129	YVCPFDGCNKFAQSTNLKSHILTH
O15391_HUMAN	1130	FVCPFDVCNRKFAQSTNLKTHILTH
TYY1_HUMAN	1131	FQCTFEGCGKRFSLDFNLRTHVRIH
O15391_HUMAN	1132	FQCTFEGCGKRFSLDFNLRTHLRIH
Q14872_HUMAN	1133	YQCTFEGCPRTYSTAGNLRTHQKTH
GLI1_HUMAN	1134	HKCTFEGCRKSYSRLENLKTHLRSH
GLI3_HUMAN	1135	HKCTFEGCTKAYSRLLENLKTHLRSH
O60255_HUMAN	1136	HKCTFEGCSKAYSRLLENLKTHLRSH
O60254_HUMAN	1137	HKCTFEGCSKAYSRLLENLKTHLRSH
O60253_HUMAN	1138	HKCTFEGCSKAYSRLLENLKTHLRSH
O60252_HUMAN	1139	HKCTFEGCSKAYSRLLENLKTHLRSH
GLI2_HUMAN	1140	HKCTFEGCSKAYSRLLENLKTHLRSH
O95409_HUMAN	1141	FQCEFEGCDRRFANSDDRKKHMVH
Q15915_HUMAN	1142	FKCEFEGCDRRFANSDDRKKHMVH
ZIC3_HUMAN	1143	FKCEFEGCDRRFANSDDRKKHMVH
GLI1_HUMAN	1144	YMCEHEGCSKAFSNASDRAKHQNRTH

O60255_HUMAN	1145	YVCEHEGCNKAFSNASDRAKHQRTH
O60254_HUMAN	1146	YVCEHEGCNKAFSNASDRAKHQRTH
O60253_HUMAN	1147	YVCEHEGCNKAFSNASDRAKHQRTH
O60252_HUMAN	1148	YVCEHEGCNKAFSNASDRAKHQRTH
GLI3_HUMAN	1149	YVCEHEGCNKAFSNASDRAKHQRTH
GLI2_HUMAN	1150	YVCEHEGCNKAFSNASDRAKHQRTH
Z143_HUMAN	1151	YVCTVPGCDKRFTEYSSLYKHHVVH
TF3A_HUMAN	1152	FKCTQEGCGKHFASPSKLKRHAKAH
TF3A_HUMAN	1153	FVCDYEGCGKAFIRDYHLSRHILTH
Q14872_HUMAN	1154	FECDVQGCEKAFNTLYRLKAHQRLH
Q14872_HUMAN	1155	FVCNQEKGKAFLTSHSLRIHVRVH
ZN76_HUMAN	1156	YRCDFPSCGKAFATGYGLKSHVRTH
Z143_HUMAN	1157	YQCEHAGCGKAFATGYGLKSHVRTH
Q14872_HUMAN	1158	FRCDHDGCGKAFAAASHHLKTHVRTH
O00153_HUMAN	1159	FICPAEGCGKSFYVLQRLKVHMRTH
ZN76_HUMAN	1160	FQCPFEGCGRSFTTSNIRKVHVRTH
Z143_HUMAN	1161	FKCPFEGCGRSFTTSNIRKVHVRTH
Q15915_HUMAN	1162	FPCPFPGCGKVFARSENLKIHKRTH
O95409_HUMAN	1163	FPCPFPGCGKVFARSENLKIHKRTH
ZIC3_HUMAN	1164	FPCPFPGCGKIFARSENLKIHKRTH
ZN76_HUMAN	1165	YTCPEPHCGRGFTSATNYKNHVRIH
Z143_HUMAN	1166	YYCTEPGCGRAFASATNYKNHVRIH
O00153_HUMAN	1167	FMCHESGCGKQFTTAGNLKNHRRIH
ZN76_HUMAN	1168	YKCPEELCSKAKTSGDLQKHVRTH
Z143_HUMAN	1169	YRCSEDNCTSKTSGDLQKHIRTH
Q14872_HUMAN	1170	FNCESEGC SKYFTTLSDLRKHIRTH
ZN76_HUMAN	1171	FRCGYKGCGRLYTTAHLKVHERAH
Z143_HUMAN	1172	FRCEYDGC GKLYTTAHLKVHERSH
BTE1_HUMAN	1173	HKCPYSGCGKVYKGSSHLKAHYRVH
BTE2_HUMAN	1174	HYCDYPGCTKVYTKSSHLKAHLRTH
O43839_HUMAN	1175	HRCHFNGCRKVYTKSSHLKAHQRTTH
UKLF_HUMAN	1176	HRCQFNGCRKVYTKSSHLKAHQRTIH
O95600_HUMAN	1177	HQCDFAGCSKVYTKSSHLKAHLRTH
Q13118_HUMAN	1178	HICSHPGCGKTYFKSSHLKAHTRTH
O75411_HUMAN	1179	HICSHPGCGKTYFKSSHLKAHTRTH
EZF_HUMAN	1180	HTCDYAGCGKTYTKSSHLKAHLRTH
O14901_HUMAN	1181	YVCSFPGCRKTYFKSSHLKAHLRTH
SP4_HUMAN	1182	HICHIEGCGKVYGKTSHLRAHLRWH
O60402_HUMAN	1183	HICHIEGCGKVYGKTSHLRAHLRWH
EKLF_HUMAN	1184	HTCAHPGCGKS YTKSSHLKAHLRTH
WT1_HUMAN	1185	FMCAYPGCNKRYFKLSHLQMHSRKH
Q16256_HUMAN	1186	FMCAYPGCNKRYFKLSHLQMHSRKH
Q15881_HUMAN	1187	FMCAYPGCNKRYFKLSHLQMHSRKH
SP2_HUMAN	1188	HVCHIPDCGKTFRKTSSLRAHVRLH
O43167_HUMAN	1189	YACKDCHRKFMDVSQLKKHLRTH
O75467_HUMAN	1190	YACRACSKVFKVSSDLLKHLRTH
ZEP1_HUMAN	1191	YICEYCNRACAKPSVLLKHLRSH
Q02646_HUMAN	1192	YICPYCSRACAKPSVLLKHLRSH

O75362_HUMAN	1193	YACSYCGKFFRSNYYLNIHLRTH
Q92981_HUMAN	1194	YKCVQPDGKAFVSRYKLMRHMAH
O76019_HUMAN	1195	YKCVQPDGKAFVSRYKLMRHMAH
RRE1_HUMAN	1196	YACSVCNKRFWSLQDLTRHMRSH
O75626_HUMAN	1197	HECQVCHKRFSTS NLKTHLRLH
Z202_HUMAN	1198	HDCSVCVGKSFTCN SHLVRHLRTH
O75123_HUMAN	1199	YACDICGKTFTFNSDLV RHRISH
Z151_HUMAN	1200	HKCSVCSKA FVN VGDLSKHI IIH
SNAI_HUMAN	1201	YACVCGTCGKAFSRPWL LQGHVRTH
O43623_HUMAN	1202	YACVCKICGKAFSRPWL LQGHIRTH
O95409_HUMAN	1203	HVCFWEECPREGKPFKAKYKLV NHIRVH
ZIC3_HUMAN	1204	HVCYWE ECPREGKSFKAKYKLV NHIRVH
O00146_HUMAN	1205	HECKLCGASFR TKGS LIRHHRRH
O00146_HUMAN	1206	HVCQFC SRGFREKGSLVRHVRHH
IKAR_HUMAN	1207	FQCNQCGASFTQKG NLLRHI K LH
CTCF_HUMAN	1208	HKCHLCGRAFRTVTLLRNHLNTH
HKR3_HUMAN	1209	HVCEFC SHAFTQKANL NMHLRTH
Q15552_HUMAN	1210	HVCEHCNAAFRTNYHLQRHVF IH
O43591_HUMAN	1211	HVCEHCNAAFRTNYHLQRHVF IH
PLZF_HUMAN	1212	YICSECNRTFPSHTALKRHLRSH
Z151_HUMAN	1213	YVCIH CQRQFADPGALQRHVR IH
MAZ_HUMAN	1214	YICALCAKEF KNGYNL RRHEAIH
O14753_HUMAN	1215	HLCTGCGKG FNDT FDLKRHVRTH
O95365_HUMAN	1216	YECNI CKVRFTRQDKLK VHM RKH
O15156_HUMAN	1217	YACEVCGVRFTRNDKLKIHM RKH
O75066_HUMAN	1218	YSCEECGAKFAANSTLKNH LRLH
O95365_HUMAN	1219	YLCQQCGAAFAHNYDLKNHMRVH
O15156_HUMAN	1220	YSCP HCPARFLHSYDLKNHMHLH
Z151_HUMAN	1221	HKCEDCGKEFTHTGNFKRHIRIH
Z151_HUMAN	1222	YRCEDCGKLFTTSGNLKRHQLVH
Z151_HUMAN	1223	YKCRECGKQFTTSGNLKRHRLIH
O15090_HUMAN	1224	YDCPYCGKTFR TSHHLKVHLRIH

Example 3: Non-human zinc finger databases.

For providing novel combinations of non-antigenic, optimised zinc fingers, for use in species other than humans, separate species-specific zinc finger databases are required, 5 such as mouse, chicken, pig, cow, *etc.*

The fingers listed below are in a format that can be linked with classical wild-type canonical “TGEKP” linkers (i.e. ...TGEKP – zinc finger peptide sequence – TGEKP – zinc finger peptide sequence – TGEKP - etc...). For each peptide sequence, an 10 oligonucleotide is designed to encode the peptide sequence; the oligonucleotide can then be linked into a library selection system, as described in the Examples *infra*.

Mouse Zinc Finger Database.

15	544 zinc finger units		
	Name	SEQ ID NO	Peptide sequence
	O35745_MOUSE	1225	HQCTHCEKTFNRKDHLKNHLQTH
	ZFX2_MOUSE	1226	HRCEYCKKGFRRPSEKNQHIMRH
	ZFX1_MOUSE	1227	HRCEYCKKGFRRPSEKNQHIMRH
	ZFY2_MOUSE	1228	HKCDMCSKGFHRPSELKKHVATH
	ZFY1_MOUSE	1229	HKCDMCSKGFHRPSELKKHVATH
	ZFX2_MOUSE	1230	HKCDMCDKGFHRPSELKKHVAAH
	ZFX1_MOUSE	1231	HKCDMCDKGFHRPSELKKHVAAH
	ZFA_MOUSE	1232	HKCDMCDKGFHRPSELKKHVAAH
	Q9Z162_MOUSE	1233	YTCSVCGKGFSRPDHLSCHVKHVH
	MAZ_MOUSE	1234	YNC SHCGKSF SRPDHLSH VRQVH
	Q08376_MOUSE	1235	YSCEVCGKSFIRAPDLKKHERVH
	Z151_MOUSE	1236	HKCPHCDKKFNQVGNLKAHLKIH
	ZFX2_MOUSE	1237	FRCKRCKGFRQQSELKKHMKT
	ZFX1_MOUSE	1238	FRCKRCKGFRQQSELKKHMKT
	Q62518_MOUSE	1239	YVCTMCGKGYTLNSNLQVHLRVH
	Q60636_MOUSE	1240	YECSVCAKTFGQLSNLKVHLRVH
	Q9Z117_MOUSE	1241	CSCPECGV LQSHLRSHYRLH
	Q61898_MOUSE	1242	CSCPEC GREFHQLSHLRKHYRLH
	O88631_MOUSE	1243	YSCQYCGKVFHQLSHFKSHFTLH
	Q61164_MOUSE	1244	HKCPDCDMAFVTSGELVRHRRYKH
	O35483_MOUSE	1245	FRCADCGRGFAQRSNLAKHRRGH
	O35483_MOUSE	1246	FVCGVCGAGFSRRAHLT A HGRAH
	O70162_MOUSE	1247	FVCRDCGQGFVRSARLEEHRRVH
	Q9Z1D8_MOUSE	1248	HRCGDCGKFFLQASNFIQHRRIH
	O35483_MOUSE	1249	HRCPDCGKGF GHSSDFKRHRRTH
	O35483_MOUSE	1250	---ADCGKSFVYGSHLARHRRTH

O35483_MOUSE	1251	FPCPDCGKRFVYKSHLVTHRRIH
O88282_MOUSE	1252	YKCQLCRSAFRYKGNLASHRTVH
Q61065_MOUSE	1253	YKCDRCQASFRYKGNLASHKTVH
BCL6_MOUSE	1254	YKCDRCQASFRYKGNLASHKTVH
O70162_MOUSE	1255	FACQDCGRRFNQSTKLIQHQRVH
O70162_MOUSE	1256	--CVECGERFGRRSVLLQHRRVH
Q9Z0G7_MOUSE	1257	-DCPVCNKKFKMKHHLTEHMKTH
Q08376_MOUSE	1258	---HMCDKAFHKSHLKDERRH
Q64318_MOUSE	1259	HECGICRKAFKHKKHLIEHMR LH
Q64318_MOUSE	1260	FKCTECGKAFKYKHHLKEHLRIH
Q9Z1D8_MOUSE	1261	FKCNECGKGFGGRRSHLAGHLRLH
Q9Z1D8_MOUSE	1262	YGCNECGKSFGRHSHLIEHLKRH
Q9Z2X6_MOUSE	1263	---YVCKQCGKAFTLSSSLRRH
KID1_MOUSE	1264	YVCKECGKAFTLSTS LLYKHLRTH
Q9Z1D7_MOUSE	1265	HGCDEC GKSFTQHSRLIEHKRVH
ZF90_MOUSE	1266	YRCNL CGRSFRHSTS LTQHEVTH
Q9Z2X6_MOUSE	1267	YVCKECGKAFARSTSLHIHEGTH
Q9Z2X6_MOUSE	1268	YVCKHCGKAYTTYNTLRAHERSH
Q9Z2X6_MOUSE	1269	YVCKHCGKAYTTYNTLRAHERSH
Q9Z2X6_MOUSE	1270	YVCKHCGKAYTSYSTLRAHERSH
Q9Z2X6_MOUSE	1271	YVCKHCGKAYTSYSTLRAHERSH
Q9Z2X6_MOUSE	1272	YVCKHCGKAYTSYSTLRAHERSH
Q9Z2X6_MOUSE	1273	YVCKHCGKAFTQSSYLRHKRTH
ZF37_MOUSE	1274	YECEQCGKAHGKHALTDHLRIH
Q62514_MOUSE	1275	YECEQCGKAHGKHALTDHLRIH
Q61491_MOUSE	1276	YECNQCGKAFTQFFPLKRHEITH
ZF37_MOUSE	1277	YKCDEC GKA FGHSSSLTYHMRTH
Q62514_MOUSE	1278	YKCDEC GKA FGHSSSLTYHMRTH
Q61491_MOUSE	1279	YQC NQCAKAFPYHRTLQIHERTH
Q61491_MOUSE	1280	CEYNQCWKAFA YHKT LQIHERTH
Q61491_MOUSE	1281	YECNQCGKAFAC YQSFQIHKRTH
Q61491_MOUSE	1282	YECNQCGKAFAC NRYLQIHKRTH
Q61491_MOUSE	1283	YECNQCGKAFAC PRYLQIHKRTH
Q61491_MOUSE	1284	YECNQCGKAFAC LRLQNHKTTH
Q61491_MOUSE	1285	FECNQCGKAFAHHSTLQRHKRTH
Q61491_MOUSE	1286	YECNQCGKAFTRHSTLQIHKRTH
Q61491_MOUSE	1287	YECNQCGKAFTCRSNLQIHKRTH
Q9Z2X6_MOUSE	1288	YVCKQCGKAFTRSSHLQIHKITH
Q9Z2X6_MOUSE	1289	YICKQCGKAFARSSH LQIHKRSH
Q61491_MOUSE	1290	YKCKQCGKDFTHHSTLHIHKRIH
Q9Z2X6_MOUSE	1291	YSCKLCGKAFTHSNYLQIHKRIH
Q61491_MOUSE	1292	YECNQCGKAFARNSNLLDHKRIH
Q64247_MOUSE	1293	YICKQCGKTFRYLSCFQKHERIH
Q9Z2X6_MOUSE	1294	YACKQCDKAFKYLSSLQNHKRIH
Q9Z2X6_MOUSE	1295	HACKQCGKSFKRQSNVQAHERNH
Q64247_MOUSE	1296	YTCKHCTKTFTTSSTRNSHEKTH
Q64247_MOUSE	1297	YACKHCGKAFTTSSARN SHERIH
Q64247_MOUSE	1298	YACKHCGKAFTSSDRNSHERIH

Q64247_MOUSE	1299	YPCKYCGKAFATS SDRNSHERIH
Q64247_MOUSE	1300	YSCTHCGKAFSSPSDYNSCERIH
O88412_MOUSE	1301	YVCNECGKAFTCSSYLLIHQRIH
ZF35_MOUSE	1302	YMCNHCYKHFSQSSDLIKHQRIH
Q9Z2X6_MOUSE	1303	YVCKQCGKAFAQSSYLHIHQRSH
ZF38_MOUSE	1304	YQCKDCGKAFSGKGSLIRHYRIH
OZF_MOUSE	1305	YECNKCGKAFSRITSLIVHVRIH
Q9Z0Q5_MOUSE	1306	YECNECGKAFSQRTSLIVHVRIH
ZF90_MOUSE	1307	YQCNVCGKAFKRSTS FIEHHRIH
OZF_MOUSE	1308	YECKICGKAFQCQSSLTVMRSH
Q9Z0Q5_MOUSE	1309	YECNVCGKAFSQSSLTVMRSH
ZF90_MOUSE	1310	YECIDCGKAFSQSSLI QHERTH
Z151_MOUSE	1311	CQCVICGKAFTQASSLIAHVRQH
OZF_MOUSE	1312	YECKGCGKAFIQKSSLIRHQRSH
Q9Z0Q5_MOUSE	1313	FECKDCGKAFIQKSNLIRHQRTH
Q9Z162_MOUSE	1314	---TYCSKA FRD SYHLRRHQ SCH
Q9Z162_MOUSE	1315	HACEMCGKA FRD VYHLNRHKL SH
MAZ_MOUSE	1316	HACEMCGKA FRD VYHLNRHKL SH
Q61898_MOUSE	1317	FRCTECDKS FIRS SHLREHQ KIH
Q60585_MOUSE	1318	FDCKECGKTFSRGYHLTLHQRIH
O35483_MOUSE	1319	YACAEGR RFGQSA ALTRHQ WAH
Q60585_MOUSE	1320	YACTECGKSFRQVAH LTRHQ RL N
Q9Z1D9_MOUSE	1321	YACPECGE CFRQSSHLSRHQR TH
Q9Z1D9_MOUSE	1322	YKCFQCGERFRQSTHLVRHQR IH
O88631_MOUSE	1323	YKCTKCDKLFTQYSHLRRHQR Y
Q60585_MOUSE	1324	YKCTECKKA FRQHSHLTYHQ RIH
MLZ4_MOUSE	1325	HKCTECAKASAASPHL IQHQ RT H
Q9Z116_MOUSE	1326	YECTECSKA FCQKSHL TQHQ RVH
O70237_MOUSE	1327	YPCQFCGKRF HQKSDM KKHTYIH
GFI1_MOUSE	1328	YPCQYCGKRF HQKSDM KKHTFIH
Q61624_MOUSE	1329	FRCDEC GMRF I QKYH MERH KR TH
P97475_MOUSE	1330	FRCDEC GMRF I QKYH MERH KR TH
Q61624_MOUSE	1331	FQCSQCDMRF I QKYLLQRHEKIH
P97475_MOUSE	1332	FQCSQCDMRF I QKYLLQRHEKIH
ZFP1_MOUSE	1333	FVCNYCDKTFSFKSLLVSHKRIH
Q9Z116_MOUSE	1334	YICFECRKA FYRK SELTDHQ RIH
Q9Z116_MOUSE	1335	YECKECGKA FCQK PQLTLHQ RIH
ZFP1_MOUSE	1336	YGCSECGKTFA QKFELT THQ RIH
Q06054_MOUSE	1337	YKCSDCGKCF I QKANLR THQ KIH
Q06054_MOUSE	1338	YKCSDCGKCF I QKANLR THERI H
Q06054_MOUSE	1339	YKCSDCDKCF I QKAKL KKHQ RIH
Q06054_MOUSE	1340	YKCSECDKCF I QKDHLR THQ RL H
Q06054_MOUSE	1341	YKCSECDKCF I FIRKANL RRH RHI H
Q06054_MOUSE	1342	YKCSECHKCF I FIRKA HL RRH QRIH
Q06054_MOUSE	1343	YKCSECHKCF I QQAHL RRH QKIH
Q06054_MOUSE	1344	YI CAECN KCF I QKSQLK THQ RIH
MLZ4_MOUSE	1345	HIC SQCGKA FSQI S D LNRH QK TH
ZF37_MOUSE	1346	YECNECGIA FSQK SHL VVHQ RT H

Q62514_MOUSE	1347	YECNECGIAFSQKSHLVLHQRT
ZF37_MOUSE	1348	YECVECGKAFSQKSHLIVHQRP
Q62514_MOUSE	1349	YECVECGKAFSQKSHLIVHQRT
ZF37_MOUSE	1350	FECNECGKTFSKKSHLVIHQRT
Q62514_MOUSE	1351	FECNECGKTFSKKSHLVIHQRT
MFG3_MOUSE	1352	FECKECGKAFHFSSQLNNHKTSH
Q62514_MOUSE	1353	FECYECGKAFNAKSQLVIHQRS
ZF37_MOUSE	1354	FECYECGKAFNAKSQLVIHQRS
Q9Z116_MOUSE	1355	YECKICGKCFYWKTFSNRHQSTH
O88412_MOUSE	1356	YSCNECGKAFRQKSSLTVHQRT
Q9Z116_MOUSE	1357	YECAECGKAFSTKSYLTVHQRT
P70405_MOUSE	1358	YECSKCGKTFRGKYSLDQHQRV
ZF90_MOUSE	1359	HECADCGKTFLWRTQLTEHQRI
KR2_MOUSE	1360	YECMICGKHFTGRSSLTVHQVI
KR2_MOUSE	1361	YECDQCGKAFIKNSSLIVHQRI
Q9Z1D7_MOUSE	1362	YKCSVCGKAFIQKISLIEHEQI
Q61116_MOUSE	1363	YKCDTCGKAFSQKSSLQVHQRI
O70237_MOUSE	1364	--CRMCGKAFKRSSTLSTHLLI
GFI1_MOUSE	1365	-DCKICGKSFKRSSTLSTHLLI
Q9Z150_MOUSE	1366	HSCGICGKCFQKSTLHDHNLH
Q9Z1D7_MOUSE	1367	YKCEVCGKTFRWRTVLIRHKVV
ZF35_MOUSE	1368	-YKCMCGKAFSQCSAFTLHQRI
ZF38_MOUSE	1369	YKCKECGKAFNHSSNFNKHRI
OZF_MOUSE	1370	YGCNECGKAFSQFSTLALHMR
Q9Z0Q5_MOUSE	1371	YGCNECGKAFSQFSTLALHLRI
ZFP1_MOUSE	1372	YECTECGKTFQRSTLRLHLRI
MLZ4_MOUSE	1373	YKCDECCKNFSQNSDLVRHRR
Q62514_MOUSE	1374	YECNECGKAFKYGSSLTKHMRI
ZF37_MOUSE	1375	YECNECGKAFKYGSSLTKHMRI
KR2_MOUSE	1376	YKCHDCGKAFSKNSSLTQHRR
P70405_MOUSE	1377	CRDCGKFFSQTSHLNDHRRHTG
Q61117_MOUSE	1378	YKCSTCGKGFSRSSDLNVHCR
ZF92_MOUSE	1379	YLCQQCGKSFSRSFNLIKHR
ZF29_MOUSE	1380	YACKECGESFSYNSNLIRHQRI
O88282_MOUSE	1381	YRCSICGARFNRPANLKTHSR
Q61065_MOUSE	1382	YRCNICGAQFNRPANLKTHTR
BCL6_MOUSE	1383	YRCNICGAQFNRPANLKTHTR
ZF29_MOUSE	1384	YKCRDCGKSFSRSANLITHQRI
Q9Z1D7_MOUSE	1385	YQCLQCNKSFNRRSTLSHQGV
ZF35_MOUSE	1386	YPCNSCSKSFSSRGSDLIKHQ
ZF35_MOUSE	1387	YPCSWCIKSFSRSSLIKHQ
ZF35_MOUSE	1388	YPCNQCTKSFSSRLSDLINHQ
ZFP1_MOUSE	1389	YECDVCKTFSHKANLIKHQRI
ZF35_MOUSE	1390	YECDKCGKTFSQSSNLILHQRI
O88412_MOUSE	1391	YECNECGKTFTRSSNLIVHQRI
MLZ4_MOUSE	1392	YDCNECGKSFGRSSHЛИQHQT
MLZ4_MOUSE	1393	YECTACGKSFSRSSHLITHQK
KR2_MOUSE	1394	YECTECGKAFSQSAYLIEHRR

ZF90_MOUSE	1395	YACKECGRNFSRSSALTKHHRVH
MLZ4_MOUSE	1396	YECTECDKSFSRSSALIKHKRVH
P70405_MOUSE	1397	YKCSECGKSFSQSSILIQHRRIH
P70405_MOUSE	1398	YKCSECGNSFSQSAILNQHRRIH
Q9Z1D8_MOUSE	1399	HQCNECGKSFIQSAHLIQHRRIH
KID1_MOUSE	1400	YRCQECGMSFGQSSALIQHRRIH
P70405_MOUSE	1401	YECSQCGKSFSQKSGLIQHQVVH
P70405_MOUSE	1402	YECRECGKSFSQLATLIKQRVH
P70405_MOUSE	1403	YECSQCGKSFSQKATLVKKRVRH
Q9Z1D8_MOUSE	1404	HQCNECGRGFSLKSHLSQHQRIH
OZF_MOUSE	1405	YQCSECGKAFSQKSHHIRHQRIH
Q9Z0Q5_MOUSE	1406	YQCSECGKAFSQKSHHIRHQKIH
O88412_MOUSE	1407	YDCSECGKAFSQLSCLIVHQRIH
ZF35_MOUSE	1408	YKCSECGKAFNQSSVLILHQRIH
ZF35_MOUSE	1409	YKCDVCGKAFSQSSDRILHQRIH
KID1_MOUSE	1410	FKCNTCGKTFRQSSSRIAHQRIH
OZF_MOUSE	1411	YKCNECGTIFRQKQYLIKHHNIH
Q9Z0Q5_MOUSE	1412	FKCNECGTAFGQKKYLIKHQNIH
OZF_MOUSE	1413	FECSQCGRAFSQKQYLIKHQNIH
Q9Z0Q5_MOUSE	1414	FECNECGKAFSQKQYVIKHQSTH
OZF_MOUSE	1415	FKCNECGKAFSQKENLIIHQRIH
Q9Z0Q5_MOUSE	1416	FECSDCGKAFSQKENLLTHQKIH
KID1_MOUSE	1417	FKCSECGRAFSQASLIQHERIH
O88412_MOUSE	1418	FECHECGKAFIQSANLVVHQRIH
O88412_MOUSE	1419	FTCSECGKGFSQSANLVVHQRIH
O88412_MOUSE	1420	FACSDCGKAFTQSANLIVHQRSW
KR2_MOUSE	1421	YKCHECGKAFSQSMNLTVHQRTW
ZF38_MOUSE	1422	YQCNECGKSFSQHAGLSSHQRLH
KID1_MOUSE	1423	YNCNECGKALSSHSTLIIHERIH
O35700_MOUSE	1424	YKCDQCPKAFNWKSNLIRHQMSH
EVI1_MOUSE	1425	YKCDQCPKAFNWKSNLIRHQMSH
Q62518_MOUSE	1426	YKCDVCGKSFGRWSNLIIHHRIH
Q9Z1D8_MOUSE	1427	YACHLCGKAFRVRSHLVQHQSVH
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Q9Z1D7_MOUSE	1429	YECNDCGKAFVYNSSLATHQETH
MFG3_MOUSE	1430	YKCNACGRAFNRRSNLMQHEKIH
MFG3_MOUSE	1431	YKCNVCGKAFNRRSNLLQHQKIH
O88412_MOUSE	1432	YVCKGKCGKAFTQSSNLTVHQKIH
Q9Z116_MOUSE	1433	YECKECKRKAFYDKSNLKRHQKIH
Q60585_MOUSE	1434	YECKECKRKFRRYSELISHQGIH
Q60585_MOUSE	1435	YECKECKGKAFRQCAHLSRHQRIH
ZF37_MOUSE	1436	YECIECGKAFKQNASLTKHMKIH
Q62514_MOUSE	1437	YECIECGKAFKQNASLTKHMKIH
Q61849_MOUSE	1438	YECNECGKAFKRHSFVRHQKIH
MFG3_MOUSE	1439	FECKDCGKVFRNLNIHLIRHQRFH
Q61849_MOUSE	1440	YECKECKGKAFRLPQQLTRHQKCH
Q06054_MOUSE	1441	HRCNECGKSLSSSSGLQRHQRIH
O35700_MOUSE	1442	HACPECGKTFATSSGLQHKHIIH

EVI1_MOUSE	1443	HACPECGKTFATSSGLKQHKHIH
ZF92_MOUSE	1444	YECGECGKTFTRSSNLVKHQVIH
O88412_MOUSE	1445	FKCSECEKAFSYSSQLARHQKVH
ZF90_MOUSE	1446	FECNVCGKAFRHSSLGQHENAH
KID1_MOUSE	1447	YECNTCGKLFNHRSSLTNHYKIH
ZF29_MOUSE	1448	YKCDECCKGSFSDGSNFSRHQTTH
OZF_MOUSE	1449	YKCGECGKAFSQRGNFLSHQKQH
O70162_MOUSE	1450	CDVCGKVFSQRSNLLRHQKIHTG
ZFP1_MOUSE	1451	YECNECAKTFKKSNLIIHQKIH
O88412_MOUSE	1452	YKCKDCEKAFSCFSHLIVHQRIH
Q9Z1D7_MOUSE	1453	YKCNECGRAFGQWSALNQHQRLH
ZF90_MOUSE	1454	YQCSLCGKAFQRSSLVQHQRIH
Q64247_MOUSE	1455	-----CGKVFILSGDLIKHERIH
MFG3_MOUSE	1456	YECEQCGSAFRRLPYQLTQHQRIH
Q61849_MOUSE	1457	FECELCGSAFRCRSQLNKLRIH
MFG3_MOUSE	1458	FKCKLCESAFRRKYQLSEHQRIH
Q61849_MOUSE	1459	FKCQECGKAFVVLAYLIEHQSIH
Q64247_MOUSE	1460	FVCKQCGEAFVNSSHLSHERIH
MFG3_MOUSE	1461	FQCKECGKAFSRCSSLVQHERTH
Q64247_MOUSE	1462	FVCKTCGKAFSRSDYLINHKRIH
ZF90_MOUSE	1463	FVCKKCGKAFKRLGHFMNHERIH
MFG3_MOUSE	1464	FQCKECGKAFSRCSSLVQHERTH
MFG3_MOUSE	1465	FECKDCGKAFTVLAQLTRHQTIH
MFG3_MOUSE	1466	FHCKVCGKAFTVLAQLTRHENIH
MFG3_MOUSE	1467	FECKECGKSFKRVSSLVEHRIIH
ZFP1_MOUSE	1468	FECPECGKAFTHQSNLIVHQRAH
ZF92_MOUSE	1469	FECTECGKAFSRSSNLIEHQRIH
O54978_MOUSE	1470	FECQECGGEAFARSELIEHQKIH
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O70162_MOUSE	1472	FACAECGQSFRQRSNLQHQRIH
O70162_MOUSE	1473	FACPECGQSFRQHANLTQHRRIH
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O70162_MOUSE	1475	AECGKTFRQRATLTQHLCVHTGE
Q9Z1D8_MOUSE	1476	FRCEECGKSYNQRVHЛИQHHRVH
Q9Z1D8_MOUSE	1477	FKCGECGKSYNQRVHLTQHQRVH
ZF37_MOUSE	1478	FECNQCGKAFKQIEGLTQHQRVH
Q62514_MOUSE	1479	FECNQCGKAFKQIEGLTQHQRVH
O88282_MOUSE	1480	YPCPTCGTRFRHLQTLKSHVRIH
Q61065_MOUSE	1481	YPCEICGTRFRHLQTLKSHLRIH
BCL6_MOUSE	1482	YPCEICGTRFRHLQTLKSHLRIH
Q60585_MOUSE	1483	YDCKECGKAFRVRQQLTLHERIH
Q60585_MOUSE	1484	YDCKECGKAFRVRGQQLMLHQRIH
Q60585_MOUSE	1485	YECGECGKAFKVRQQLTFHQRIH
OZF_MOUSE	1486	YACKECGKAFNGKSYLNKEHEKIH
OZF_MOUSE	1487	YTCKECGKAFSGKSNLTEHEKIH
Q9Z0Q5_MOUSE	1488	FICKECGKTFSGKSNLTEHEKIH
MFG3_MOUSE	1489	YKCKDCGKCFGCKSNLHQHESIH
Q61849_MOUSE	1490	YQCKECGKCFRORSKLTEHESIH

Q61849_MOUSE	1491	YECKECGKCFGCRSTLTQHQSvh
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ZF92_MOUSE	1493	FVCRMCGKVRRSFALLEHTRIH
ZF92_MOUSE	1494	YECSECGKQFQRSLALLEHQRIH
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P70405_MOUSE	1496	YECSECGLFRQNSSLVDHQKTH
REX1_MOUSE	1497	HVCAECGKAFTESSKLKRHFLVH
TYY1_MOUSE	1498	HVCAECGKAFTESSKLKRHQLVH
ZFX2_MOUSE	1499	HICVECGKGFRHPSELKKHMRIH
ZFX1_MOUSE	1500	HICVECGKGFRHPSELKKHMRIH
ZFA_MOUSE	1501	HICVECGKGFCHPSELKKHMRIH
ZFY2_MOUSE	1502	HICGECKGKGFRHPSALKKHIRVH
ZFY1_MOUSE	1503	FICGECKGKGFRHPSALKKHIRVH
Q61116_MOUSE	1504	--CHECGKGFRQSSALQTHQRVH
Q06054_MOUSE	1505	YQCRKCGKCFRTYSSLYRHRRTH
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KR2_MOUSE	1511	YQCKECGKAFRKNSSLIQHERIH
KID1_MOUSE	1512	YLCNECGNTFKSSSSLRYHQRIH
KR2_MOUSE	1513	YGCDECGBTFRQSSLLKHQRIH
ZF37_MOUSE	1514	YKCNECGKTFRHSSNLMQHLRSH
Q62514_MOUSE	1515	YKCNECGKTFRHSSNLMQHLRSH
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Q62514_MOUSE	1518	YECKECGKSFRYNSSLTEHVRTH
Q9Z117_MOUSE	1519	YKCKECGKSFLELSHLKRHYRIH
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Q61898_MOUSE	1521	YECKECGKSFIELSHLKTHYRIH
Q9Z1D7_MOUSE	1522	HGCDECGBTQHSRLIEHKRVH
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O89090_MOUSE	1524	--CPECPKRFMRSDHLSKHIKTH
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O89087_MOUSE	1526	--CPECPKRFMRSDHLSKHIKTH
Q62445_MOUSE	1527	--CPECSKRFMRSDHLSKHKVTH
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Q61596_MOUSE	1529	--CPMCDRRFMRSDHLLTKHARRH
BTE1_MOUSE	1530	--CPLCEKRFMRSDHLLTKHARRH
Q62445_MOUSE	1531	FICNWMFCGKRFTRSDELQRHRRTH
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EZF_MOUSE	1536	YHCDWDGCGWKFARSDELTRHYRKH
Q60980_MOUSE	1537	YKCTWEGCTWKFARSDELTRHFRKH
O35738_MOUSE	1538	YKCTWEGCTWKFGRSDELTRHYRKH

Q9Z0Z7_MOUSE	1539	YKCTWEGCDWRFARSDELTRHYRKH
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EKLF_MOUSE	1541	YACSWDGCWDWRFARSDELTRHYRKH
Q61596_MOUSE	1542	FSCSWKGCCRFFARSDELSRHRTH
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BTE1_MOUSE	1544	FPCTWPDCLKKFSRSDELTRHYRTH
EGR2_MOUSE	1545	YPCPAEGCDRRFSRSDELTRHIRIH
WT1_MOUSE	1546	YQCDFKDCERRFSRSRSDQLKRHQRRH
WT1_MOUSE	1547	FQCKTCQRKFSRSRSDHLKTHTRTH
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KR2_MOUSE	1549	YQCNECGKPFSRSTNLTRHQRTH
O35700_MOUSE	1550	YTCRYCGKIFPRSANLTRHLRTH
EVI1_MOUSE	1551	YTCRYCGKIFPRSANLTRHLRTH
ZF29_MOUSE	1552	FQCAECGKSFSRSPNLIAHQRTH
ZF38_MOUSE	1553	YVCTKCGKAFSHSSNLTLYRTH
Q9Z1D8_MOUSE	1554	YQCDSCGKAFSYSSDLIQHYRTH
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ZF29_MOUSE	1556	YRCPECGKGFNSNSNFITHQRTH
ZF38_MOUSE	1557	YICAECKAFNSSSNLTKHRRTH
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ZF90_MOUSE	1559	YECNECGEAFSRLSSLTQHERTH
MLZ4_MOUSE	1560	YHCNECGENFSRISHLVHQRTH
ZF29_MOUSE	1561	YKCLMCGKSFSRGSILVMHQRAH
MLZ4_MOUSE	1562	YECEECGKSFSRSSHLLAQHQRTH
MLZ4_MOUSE	1563	YKCYECGKGFSRSSHLLIQHQRTH
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ZF29_MOUSE	1570	YKCTECGQKFSQSSALITHRRTH
KID1_MOUSE	1571	FKCKECSKAQFSQSSALIQHQITH
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Q9Z116_MOUSE	1577	YECTVCRKSICKSSFSHHWRTH
KR2_MOUSE	1578	YTCNVCDKHFIERSSLTVHQRTH
Q61164_MOUSE	1579	FQCSLCSYASRDTYKLKRHMRTH
P97365_MOUSE	1580	FQCWLCASAKFKISSLKRHMRVH
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ZF35_MOUSE	1582	YTCNLCSKSFSQSSDLTKHQRVH
ZF35_MOUSE	1583	YHCSSCNKAFRQSSDLILHHRVH
ZF38_MOUSE	1584	YWCSHCGKTFCSKSNLSKHQRVH
Q9Z1D9_MOUSE	1585	YKCGDCEKSFQRSDLFKHQRTH
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ZF35_MOUSE	1590	YPCAQCNCFSFSQNSDLIKHRRIH
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ZF29_MOUSE	1598	YECPOCGKTFSRKSHLITHERTH
MLZ4_MOUSE	1599	YECVQCGKGFTQSSNLITHQRVH
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ZF35_MOUSE	1604	YPCAQCCKFSQRSQSDLVNHQRVH
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Q61116_MOUSE	1606	YTCQQCGKGFSQASYFHMHQRVH
O35483_MOUSE	1607	YRCVFCGAGFGRRSYCVTHQRTH
ZF29_MOUSE	1608	YRCGDCGKGFSQRSQSQLVHVQRTH
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Z239_MOUSE	1616	YKCGECGKGFSQSSNLHIHRCTH
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ZF38_MOUSE	1618	YDCKCGKAFGQSSDLLKHQRMH
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EVI1_MOUSE	1620	YRCKYCDRSFSISSLNLQRHVRNIH
O35483_MOUSE	1621	YRCVFCGRFSFSQSSALARHQAVH
O35483_MOUSE	1622	YLCSNCGRRFQSOSHLLTHMKTH
O70162_MOUSE	1623	FVCGEGRFSFSRSSHLLRHQLTH
O88412_MOUSE	1624	YECAKCGAAFISSNHLMRHHRTH
O88631_MOUSE	1625	YKCMEDRSYIQQYSHLKRHQKVH
O88631_MOUSE	1626	YKCKECGKSYAYRTGLKRHQKIH
Z239_MOUSE	1627	YECSKCGKGFSQSSNLHIHQRTVH
Z239_MOUSE	1628	YACEECGMSFSQRSNLHIHQRTVH
MLZ4_MOUSE	1629	YECNECWRSFGERSDLIKHQRTH
MLZ4_MOUSE	1630	YECHECGRGFSERSERSDLIKHYRVH
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Q61116_MOUSE	1632	YKCGDCGKRFSCSSNLHTHQRTVH
Q62518_MOUSE	1633	YKCGECGKSFICSSNLYIHQRTVH
Q9Z150_MOUSE	1634	CPRCGKQFNHSSNLNRHMNVHRG

Q61116_MOUSE	1635	FHCSVCGKNFSRSSHFLDHQRIH
Q61116_MOUSE	1636	KCNVCQKQFSKTSNLQAHQRVH
Q62518_MOUSE	1637	YSCDVCVGKGFSSRSQLQSHQRVH
Q62518_MOUSE	1638	FKCDACGKSFSRSSSHRLSHQRVH
Q61898_MOUSE	1639	YKCRECDKSFTQRAYLRNHHNRVH
Q61898_MOUSE	1640	YKCMEDKSFTHNSNFRTHQRVH
Q9Z117_MOUSE	1641	YKCMECNKSFTQDSHLRTHQRVH
Q61898_MOUSE	1642	YKCIECDKSFTQVSHLRTHQRVH
O88631_MOUSE	1643	YKCSECDKSFTQASQLRTHQRVH
Q61898_MOUSE	1644	YKCNECDRSFTHYASLRWHQKTH
Q9Z117_MOUSE	1645	YKCKECDKSFAHCSSFRRHQKTH
Q61898_MOUSE	1646	YKCKECDKSFAHYPNFRTHQKIH
O88631_MOUSE	1647	YKCKDCDIFTNHYSSLRRHQKVH
Q9Z117_MOUSE	1648	YKCKDCDISFIQISNLRRHQKVH
Q61898_MOUSE	1649	YKCRDCDISFSQISNLRRHQKLH
Q9Z117_MOUSE	1650	FKCRECDKSFTKCSHLRRHQSVH
Q61898_MOUSE	1651	YKCRECDKSFIHSSHLLRRHQNVH
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Q06054_MOUSE	1657	YKCECGKSFTVGSDLRKHQKCH
Q61898_MOUSE	1658	YKCIECGKSFTNNSYLRTHQKVH
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Q61898_MOUSE	1660	YRCAECDKSFRCSYLRRAHQKIH
Q9Z117_MOUSE	1661	YRCKECDKSFTECSTLRAHQKIH
Q61898_MOUSE	1662	YRCKECDKSFTSCSTLKAHQSIH
Q9Z117_MOUSE	1663	YICKECGKSFTRCSYLRRAHQKIH
O88631_MOUSE	1664	YVCKECGKSLLTCAILRAHQKIH
Q61898_MOUSE	1665	YECKEKGKSFTTCSTLRIHQTIH
Q9Z117_MOUSE	1666	YICKECGKSFTKCSTLQIHQKIH
O88631_MOUSE	1667	YTCKQCGKSFTRGSTLRVHQRIH
O88631_MOUSE	1668	YKCNICDKSFTECSSLKEHRKTH
Q9Z117_MOUSE	1669	YKCEVCDKSFTVNSTLKTTHLKIH
Q61898_MOUSE	1670	YKCEICDKSFTTTTLKTTHQKIH
Q9Z117_MOUSE	1671	YKCSVCGKSFTQCTNLKTTHQRLH
Q61898_MOUSE	1672	YKCSVCDKSFTQCTHLKIHQRRH
KID1_MOUSE	1673	YRCKEKGKSFGRRSGLFIHQKVH
ZF29_MOUSE	1674	YSCPECGKSFGNRSSLNTHQGIH
Q9Z117_MOUSE	1675	YKCKEKGKSFPQLSALKSHQKIH
Q61898_MOUSE	1676	YKCKECEKSFVQLSALKSHQKLH
O88631_MOUSE	1677	YKCNDCGKSFSYLSALQSHHKRH
Q08376_MOUSE	1678	FVCEMCTKGFTTQAHLKEHLKIH
Q60636_MOUSE	1679	FKCQTCNKGFTQLAHLQKHYLVH
Q61116_MOUSE	1680	YKCEVCGKGFTQWAHLQAHERIH
O88282_MOUSE	1681	YKCETCGSRFVQVAHLRAHVLIH
Q61065_MOUSE	1682	YKCETCGARFVQVAHLRAHVLIH

BCL6_MOUSE	1683	YKCETCGARFVQVAHLRAHVLIH
O88631_MOUSE	1684	YRCEVCDKWFTLSSSLSRHQKIH
Q61116_MOUSE	1685	YRCEVCGKRFPWLSLHSHQSVH
Z239_MOUSE	1686	YKCDKCGKGFRTRSSLLVHSLH
ZF29_MOUSE	1687	YKCGLCGKSFSQSSSLIAHQGTH
Q62518_MOUSE	1688	YKCVDCGKEFSRPSSLQAHQGIH
Q61117_MOUSE	1689	YRCEECGKGFSWSSLLIHQRAH
Q61117_MOUSE	1690	YKCEECGKVFSWSSYLLKAHQRVH
Q61116_MOUSE	1691	FKCEECGKEFRWSVGLSSHQRVH
Q61117_MOUSE	1692	YKCETCGKAFSRVSILQVHQRVH
Q61116_MOUSE	1693	YKCEECGKGFSASSFQSHQRVH
Q61116_MOUSE	1694	YKCGECGKGFSHASSLQAHHSVH
Q61117_MOUSE	1695	YQCAECGRGFTVESHLQAHQRSH
Q61117_MOUSE	1696	YQCEECGRGFCRASNFLAHRGVH
Q61117_MOUSE	1697	YKCEECGKGFRTRASTLLDHQRGH
Q61117_MOUSE	1698	YVCEECGKGFSQASHLLAHQRGH
Q62518_MOUSE	1699	YNCETCGSAFSQASHLQDHQRLH
ZF29_MOUSE	1700	YRCPECGKGFSWNSVLIIHQRIH
O70162_MOUSE	1701	YCCGECDLGFTQVSRLTEHQRIH
KID1_MOUSE	1702	YRCSECKGKFTSISRLNRRIIH
TYY1_MOUSE	1703	YVCPFDGCNKKFAQSTNLKSHILTH
REX1_MOUSE	1704	YQCTFEGCGKRFSLDFNLRTHIRIH
TYY1_MOUSE	1705	FQCTFEGCGKRFSLDFNLRTHVRIH
MTF1_MOUSE	1706	YQCTFEGCPRTYSTAGNLRTHQKTH
GLI_MOUSE	1707	HKCTFEGCRKSYSRLENLKTHLRSH
GLI3_MOUSE	1708	HKCTFEGCTKAYSRLLENLKTHLRSH
ZIC2_MOUSE	1709	FQCEFEGCDRRFANSSDRKKHMHVH
ZIC1_MOUSE	1710	FKCEFEGCDRRFANSSDRKKHMHVH
ZIC3_MOUSE	1711	FKCEFEGCDRRFANSSDRKKHMHVH
ZIC4_MOUSE	1712	FRCEFEGCERRFANSSDRKKHSHVH
GLI_MOUSE	1713	YMCEQEGCSKAFTSNASDRAKHQNRTH
GLI3_MOUSE	1714	YVCEHEGCNKAFSNASDRAKHQNRTH
O70230_MOUSE	1715	YVCTVPGCDKRFTEYSSLYKHHVVH
MTF1_MOUSE	1716	FECDVQGCEKAFTNLYRLKAHQRLH
MTF1_MOUSE	1717	FVCNQEGCGKAFLTSYSLRIHVRVH
O70230_MOUSE	1718	YQCEHSGCGKAFTGYGLKSHFRTH
MTF1_MOUSE	1719	FRCDHDGCGKAFAASHHLKTHVRTH
O70230_MOUSE	1720	FKCPIEGCGRSFTTSNIRKVHIRTH
ZIC4_MOUSE	1721	FPCPFPGCGKVFARSENLKIHKRTH
ZIC2_MOUSE	1722	FPCPFPGCGKVFARSENLKIHKRTH
ZIC1_MOUSE	1723	FPCPFPGCGKVFARSENLKIHKRTH
ZIC3_MOUSE	1724	FPCPFPGCGKIFARSENLKIHKRTH
O70230_MOUSE	1725	YYCTEPGCGRAFASATNYKNHVRIH
O70230_MOUSE	1726	YRCSEDNCTKSFKTSGDLQKHIRTH
MTF1_MOUSE	1727	FNCESQGCSKYFTTLSDLRKHIRTH
O70230_MOUSE	1728	FRCKYDGCCKLYTTAHLKVHERSH
BTE1_MOUSE	1729	HKCPYSGCGKVYKGSSHLSKAHYRVH
Q9Z0Z7_MOUSE	1730	--CDYNGCTKVYTKSSHLSKAHLRTH

Q60980_MOUSE	1731	HRCDYDGCKVYTKSSHKAHRRTH
O35738_MOUSE	1732	HRCDFEGCNKVKYTKSSHKAHRRTH
Q61596_MOUSE	1733	HICSHPGVGKTYFKSSHKAHVRTH
O89091_MOUSE	1734	HICSHPGCGKTYFKSSHKAHVRTH
Q60843_MOUSE	1735	HTCSYTNCGKTYTKSSHKAHLRTH
EZF_MOUSE	1736	HTCDYAGCGKTYTKSSHKAHLRTH
Q64167_MOUSE	1737	HICHIQGCGKVKYGTSHLRAHLRWH
O89090_MOUSE	1738	HICHIQGCGKVKYGTSHLRAHLRWH
O89087_MOUSE	1739	HICHIQGCGKVKYGTSHLRAHLRWH
Q62445_MOUSE	1740	HVCHIEGCGKVKYGTSHLRAHLRWH
O70261_MOUSE	1741	HTCGHEGCGKSYTSSHKAHLRTH
EKLF_MOUSE	1742	HTCGHEGCGKSYSKSSHKAHLRTH
WT1_MOUSE	1743	FMCAYPGCNKRYFKLSHLQMHSRKH
ZEP1_MOUSE	1744	YICEYCNRACAKPSVLLKHIRSH
Q61479_MOUSE	1745	YICQYCSRCAKPSVLQKHIRSH
O55140_MOUSE	1746	YICPYCSRACAKPSVLKKHIRSH
Q60636_MOUSE	1747	HECQVCHKRFSSTSNLKTHRLH
SNAI_MOUSE	1748	CVCTTCGKAFSRPWLLQGHVRTH
P97469_MOUSE	1749	CVCKICGKAFSRPWLLQGHIRTH
ZIC2_MOUSE	1750	HVCFWEECPREGKPFKAKYKLVNHIRVH
ZIC3_MOUSE	1751	HVCYWEECPREGKSFKAKYKLVNHIRVH
Q62065_MOUSE	1752	HECKLCGASFRKGSLIRHHRRH
Q62065_MOUSE	1753	HVCQFCCSRGFREKGSLVRHVRHH
IKAR_MOUSE	1754	FQCNQCGASFTQKGNNLLRHIKLH
Q9Z2Z2_MOUSE	1755	FHCNQCGASFTQKGNNLLRHIKLH
HELI_MOUSE	1756	FHCNQCGASFTQKGNNLLRHIKLH
Q61164_MOUSE	1757	HKCHLCGRAFRVTLLRNHLNTH
Q61624_MOUSE	1758	HVCEHCNAAFRTNYHLQRHVFH
P97475_MOUSE	1759	HVCEHCNAAFRTNYHLQRHVFH
Z151_MOUSE	1760	YVCTHCQRQFADPGGLQRHVRH
Q62511_MOUSE	1761	YICEYCARAFKSSHNLAVHRCMH
MAZ_MOUSE	1762	YICALCAKEFKNGYNLRRHEAIH
O88939_MOUSE	1763	YECNICKVRFRQDKLKVKHMRKH
Q64321_MOUSE	1764	--CEVCGVRFRNDKLKIHMRCMH
P97365_MOUSE	1765	PHKCEVCGKCFSRKDKLKTHMRCH
O88939_MOUSE	1766	YLCQQCGAAFAHNYDLKNHMRVH
Q64321_MOUSE	1767	YSCPHCPARFLHSYDLKNHMHMH
Z151_MOUSE	1768	HKCEDCGKEFTHTGNFKRHIRIH
Z151_MOUSE	1769	YRCGDCGKLFITSGNLKRHQLVH
Z151_MOUSE	1770	-KCRECGKQFTTSGNLKRHLRIH

Chicken database.

5	35 finger units	SEQ ID NO	
	Q92010_CHICK	1771	YSCEVCGKSFIRAPDLKKHERVH

Q90851_CHICK	1772	YPCTICGKKFTQRGTMTRHMRSH
Q90850_CHICK	1773	YPCTICGKKFTQRGTMTRHMRSH
Q90851_CHICK	1774	--CDACGMRFTRQYRLTEHMRIH
Q90850_CHICK	1775	--CDACGMRFTRQYRLTEHMRIH
CTCF_CHICK	1776	HKCPDCDMAFVTSGELVRHRRYKH
ZKR1_CHICK	1777	-TCGDCGKGFAWASHLQRHRRVH
ZKR1_CHICK	1778	HRCGDCGKGFAWASHLQRHRRVH
ZKR1_CHICK	1779	HRCGDCGKGFWASHLERHRRVH
ZKR1_CHICK	1780	--CPDCGKSF PWASHLERHRRVH
Q92010_CHICK	1781	--CHMCDKAFKKSHLKDHERRH
O42408_CHICK	1782	HECGICKKAFKKHHHLIEHMRLH
DEFI_CHICK	1783	HECGICKKAFKKHHHLIEHMRLH
O42408_CHICK	1784	FKCTECGKAFKYKHHLKEHLRIH
DEFI_CHICK	1785	FKCTECGKAFKYKHHLKEHLRIH
O42409_CHICK	1786	YPCQYCGKRFHQKSDMKKHTYIH
O42409_CHICK	1787	FECKMCGKTFKRSSTLSTHLLIH
ZKR1_CHICK	1788	YECPECGEAFSQGSHLTKHRRSH
ZKR1_CHICK	1789	YECPECGEAFSQGSHLTKHRRSH
ZKR1_CHICK	1790	YSCPECGESYSQSSHLSLVQHRRTH
O42409_CHICK	1791	HKCQVCGKAFSQSSNLITHSRKH
O57415_CHICK	1792	YQCNICDYIAADKAALIRHLRTH
CTCF_CHICK	1793	FQCSLCSYASRDTYKLKRHMRTH
O57415_CHICK	1794	YKCQTCERTFTLKHSVLVRHQRIH
Q92010_CHICK	1795	FVCEMCTKGFTTQAHLKEHLKIH
O57415_CHICK	1796	-TCPYCPRVFWSWASSLQRHMLTH
O57415_CHICK	1797	HSCSICGKSLSSASSLDRHMLVH
O57415_CHICK	1798	--CTVCNKRFWSLQDLTRHMRSH
Q91051_CHICK	1799	CVCKICGKAFSRPWLLQGHIRTH
O12939_CHICK	1800	CVCKMCGKAFSRPWLLQGHIRTH
O57415_CHICK	1801	YKCSVCGQSFTTNGNMHRHMKIH
IKAR_CHICK	1802	FQCNQCGASFTQKGNLLRHIKLH
CTCF_CHICK	1803	HKCHLCGRAFRTVTLLRNHLNTH
O93567_CHICK	1804	YECNICNVRFTRQDKLKVHMRKH
O93567_CHICK	1805	YLCQQCGAAFAHNYDLKNHMRVH

Plant Database.

52 finger units

SEQ ID NO

O22089_PETHY	1806		HECSICGEQFLLQALGGHMRKH
O22088_PETHY	1807		HECSFCGEDFPTGQALGGHMRKH
O22087_PETHY	1808		-ECSFCGEDFPTGQALGGHMRKH
Q39092_ARATH	1809		HKCKLCKWSFANGRALGGHMRSH
Q39217_ARATH	1810		HKCSICCSQSFTGQALGGHMRHH
P93713_PETHY	1811		HECSICGLEFAIGQALGGHMRHH
O22086_PETHY	1812		HECSICGLEFPFIGQALGGHMRHH
O22085_PETHY	1813		HECSICGMFSLGQALGGHMRHH
O22084_PETHY	1814		HECSICGMFSLGQALGGHMRHH
Q42453_ARATH	1815		HPCPICGVKFPMGQALGGHMRHH
Q42410_ARATH	1816		HPCPICGVFPMQALGGHMRHH
O65150_TOBAC	1817		HVCSICHKAFTPQALGGHKRRH
Q40897_PETHY	1818		HVCSICHKAFTPQALGGHKRRH
Q40896_PETHY	1819		HVCSICHKAFTPQALGGHKRRH
Q42430_WHEAT	1820		HRCSICQKEFPTQALGGHKRKH
Q40899_PETHY	1821		HECSICHKCFPTQALGGHKRCH
P93166_SOYBN	1822		HECSICHKSFTPQALGGHKRCH
Q96289_ARATH	1823		HVCTICNKSFPSQALGGHKRCH
Q42423_ARATH	1824		HVCTICNKSFPSQALGGHKRCH
O22533_ARATH	1825		HVCSICHKSFATGQALGGHKRCH
Q40898_PETHY	1826		HECSICHKCFSSGQALGGHKRRH
Q38895_ARATH	1827		YTCSFKREFRSAQALGGHMNVH
O23621_ARATH	1828		YTCNFCRREFRSAQALGGHMNVH
O80942_ARATH	1829		YTCSFCRREFKSAQALGGHMNVH
P93714_PETHY	1830		HECSYCGAEFTSGQALGGHMRHH
Q43614_PETHY	1831		HECAICGAEFTSGQALGGHMRHH
O22083_PETHY	1832		HECSICGAEFTSGQALGGHMRHH
Q41070_PEA	1833		HECSICGAEFTSGQALGGHMRHH
Q42375_ARATH	1834		HKCNICFRVFSSGQALGGHMRCH
O65499_ARATH	1835		HECPVCFRVFSSGQALGGHKRTH
O22090_PETHY	1836		HECPVCYRVFSSGQALGGHKRSH
O22082_PETHY	1837		HECSICHRVFSTGQALGGHKRCH
P93717_PETHY	1838		HTCSICFKSFSSGQALGGHKRCH
O04177_BRARA	1839		HTCSICFKSFSSGQALGGHKRCH
O04176_BRARA	1840		HQCSICHRVFSSGQALGGHKRCH
P93715_PETHY	1841		HECPICAKVFTSGQALGGHKRSH
Q39092_ARATH	1842		HECPYCDRVFKSGQALGGHKRSH
P93719_PETHY	1843		HACPPCPRMFKSGQALGGHKRSH
P93718_PETHY	1844		YECPLCFKIFQSGQALGGHKRSH
O22091_PETHY	1845		-KCSVCGKSFSSYQALGGHKTS
Q42430_WHEAT	1846		YKCTVCGKSFSSYQALGGHKTS
O04177_BRARA	1847		YKCTVCGKSFSSYQALGGHKTS
O04176_BRARA	1848		YKCSVCDKTFSSYQALGGHKASH
Q96289_ARATH	1849		YKCSVCDKTFSSYQALGGHKASH
Q42423_ARATH	1850		YKCSVCDKSFSSYQALGGHKASH
Q40897_PETHY	1851		YKCSVCDKSFSSYQALGGHKASH
Q40896_PETHY	1852		YKCNVCNKSFHQSQALGGHKASH
Q40898_PETHY	1853		YKCSVCDKAFSSYQALGGHKASH
O65150_TOBAC	1854		YKCSVCDKSFPSYQALGGHKASH
P93166_SOYBN	1855		YKCSVCGKGFGSYQALGGHKASH
Q40899_PETHY	1856		YKCSVCDKAFTSSYQALGGHKASH
O22533_ARATH	1857		

Arabidopsis Database

SEQ ID NO

Q9ZU64/169-191	1858	YTCPKCNSIFDTSQKFAAHMSSH
O23621/40-62	1859	YTCNFCRREFRSAQALGGHMHNVH
O23504/5-27	1860	HKCKLCSKSFCNGRALGGHMKSH
Q9SYC5/250-275	1861	WYCSCGSDFHKRSLKDHVKAFCGNGH
Q9SYC5/224-246	1862	FACRMCGKAFAVKGDWRTHEKNC
O22533/89-111	1863	YKCSVCDKAFSSYQALGGHKASH
O22533/148-170	1864	HVC SIC HKSFATGQALGGHKRCH
Q9SN24/149-171	1865	HNCSICFKSFPSGQALGGHKRCH
Q9SN24/94-116	1866	YKCSVCGKSFPSYQALGGHKTSH
Q9STI7/117-140	1867	YFCGVCDRFRYTNEKLINHFKQIH
Q9STM3/1296-1320	1868	LKCPWKGCKMTFKWAWSRTEHIRVH
Q9STM3/1243-1268	1869	YQCNMEGCTMSFSSEKQLMLHKRNIC
Q9STM3/1271-1290	1870	KCGGKNNFSHKYLVQHQRVH
Q9STM3/1326-1352	1871	YVCAEPDCGQTFRFVSDFSRHKRKTGH
Q9STM3/1296-1320	1872	LKCPWKGCKMTFKWAWSRTEHIRVH
O81801/142-164	1873	PMCNVCGKGFA SWKAVFGHLRQH
O65601/61-83	1874	QKCEKCSREFCSPVNRRHNRMH
Q9SFY6/118-140	1875	YKCSVCDKTFSSYQALGGHKASH
Q9SFY6/174-196	1876	HVCTICNKSFPSGQALGGHKRCH
O65245/147-171	1877	FYCELC SKQYRTVMEFEGLSSYDH
Q39261/52-74	1878	FSCNYCQRKFYSSQALGGHQNAH
Q9SSW0/118-140	1879	HVC S VCGKS FATGQALGGHKRCH
Q9SSW0/75-97	1880	YKCGV CYKTFSSYQALGGHKASH
Q39262/61-83	1881	FSCNYCQRKFYSSQALGGHQNAH
Q9SSW1/164-186	1882	HTCSICFKSFASGQALGGHKRCH
Q9SSW1/97-119	1883	YKCTVCGKSFSSYQALGGHKTSH
Q9ZPT0/145-167	1884	WVCERCSKG YAVQSDYKAHLKTC
Q9ZPT0/67-89	1885	YICEICNQGFQRDQNLQMRRRH
Q9ZPT0/172-193	1886	HSCDCGRVFSRVESFIEHQDNC
Q39263/85-107	1887	FSCNYCQRKFYSSQALGGHQNAH
Q9SGD1/291-316	1888	WYCTCGSDFHKRSLKDHIRSGSGH
Q9SGD1/265-287	1889	FSCGKCGKALAVKGDWRTHEKNC
Q9SGD1/180-202	1890	FAC SIC SKTFNRYNNMQMHWGH
Q9SSW2/106-128	1891	YKCNVCEKAFPSYQALGGHKASH
Q9SSW2/165-187	1892	HECSIC HKVFPTGQALGGHKRCH
Q39264/60-82	1893	HECQYCGKEFANSQALGGHQNAH
P93815/7-30	1894	QECAVCKRVLSSHQLISHYNAAH
Q39265/41-63	1895	YECQYCCREFANSQALGGHQNAH
Q39266/59-81	1896	FSCNYCRRKFYSSQALGGHQNAH
Q39267/93-115	1897	FECHYCFRNFP TSQALGGHQNAH
Q9SVY1/301-323	1898	FMCRKCGKAFAVRGDWRTHEKNC
Q9SVY1/217-239	1899	FSCPVCFKTFNRYNNMQMHWGH
Q9SGH2/1804-1827	1900	IHC LICH KTFASDDEFEDHTESKC
Q38895/47-69	1901	YTCSFCKREFRSAQALGGHMHNVH
Q9SLB8/49-71	1902	YTCSFCRREFRSAQALGGHMHNVH
Q9SL35/188-210	1903	HECSICGSEFTSGQALGGHMRRH
Q9SL35/113-135	1904	YECKTCNRTFSSFQALGGHRASH

081013/49-71	1905	HFCVICEKQFSSGKAYGGHVRIH
081013/119-141	1906	IRCCCLCGKEFQTMHSLFGHMRRH
023395/664-686	1907	LHCEKCGKALQPTEMEKHLKVFH
Q9SI97/34-56	1908	FACKTCNKEFPSFQALGGHRASH
Q9SI97/78-100	1909	HECPICGAEFAVGQALGGHMRKH
Q9SR34/35-57	1910	YVCSFCIRGFSNAQALGGHMINH
Q42485/68-90	1911	FSCNYCQRKFYSSQALGGHQNAH
082389/126-149	1912	FPCNSCGEIFPKINLLENHIAIKH
Q9SQX8/182-204	1913	YQCKTCDRTFPSFQALGGHRASH
Q9SQX8/261-283	1914	HECGICGAEFTSGQALGGHMRRH
065499/222-244	1915	HKCNICFRVFSSGQALGGHMRCH
065499/77-99	1916	RPCTECGRKFWSWKALFGHMRC
065499/162-184	1917	FECCGGCKKVFGSHQALGGHRASH
Q9SCM4/220-243	1918	DVCPKCSRGRDPPV DLLKHIDKDH
Q96289/80-102	1919	YKCSVCDKTFSSYQALGGHKASH
Q96289/136-158	1920	HVCTICNKSFPSGQALGGHKRCH
Q9SCQ6/139-161	1921	WKCDKCSKKYAVQSDWKAHSKIC
Q9SCQ6/166-187	1922	YKCDCGTLFSRRDSFITHRAFC
Q9SCQ6/63-85	1923	FVCEICNKGFQRDQNLQLHRRGH
Q9SFS1/70-92	1924	YVCEICNQGFQRDQNLQMHRRRH
Q9SFS1/148-170	1925	WICERCSKGYAVQSDYKAHLKTC
Q9SFS1/175-196	1926	HSCDCGRVFSRVESFIEHQDT
Q9SSA6/575-598	1927	IHCCLICHKTFASDDEFEDHTESKC
Q42410/39-61	1928	FTCKTCLKQFHSFQALGGHRASH
Q42410/82-104	1929	HPCPICGVFPMQALGGHMRH
Q9XFP6/12-35	1930	VWCYYCDREFDDEKILVQHQKAKH
Q9XFP6/36-59	1931	FKCHVCHKKLSTASGMVIHVLQVH
O22238/218-241	1932	VSCGSCKTFNSGNALESHNKA
Q42453/40-62	1933	FRCKTCLKEFSSFQALGGHRASH
Q42453/86-108	1934	HPCPICGVKFPQALGGHMRH
Q42375/113-135	1935	YECKTCNRTFSSFQALGGHRASH
Q42375/188-210	1936	HECSICGSEFTSGQALGGHMRH
O22759/159-181	1937	WKCEKCSKFYAVQSDWKAHTKIC
O22759/186-207	1938	YRDCCGTLFSRKDTFITHRAFC
O22759/82-104	1939	FVCEICNKGFQRDQNLQLHRRGH
Q9ZUL3/81-103	1940	FICEVCNKGFQREQNLQLHRRGH
Q9ZUL3/157-179	1941	WKCDKCSKRYAVQSDWKAHSKTC
Q9ZUL3/184-205	1942	YRDCCGTLFSRRDSFITHRAFC
P93751/95-117	1943	FECHYCFRNFPPTSQALGGHQNAH
O81827/196-219	1944	VSCHKCGEKF SKLEAAEAHHTKH
Q9ZUL4/82-104	1945	WKCEKCSKRYAVQSDWKAHSKTC
Q9ZUL4/109-130	1946	YRDCCGTIFSRRDSYITHRAFC
Q9ZUL4/6-28	1947	FICDVCNKGFQREQNLQLHRRGH
Q9SHD0/194-216	1948	FKCETCGVKFKSYQALGGHRASH
Q9SHD0/243-265	1949	HECPICFRVFTSGQALGGHKRSH
Q9SHD0/4-26	1950	YKCRFCFKSFINGRALGGHMRSH
O64936/131-153	1951	YQCNVCGRELPSYQALGGHKASH

O64936/179-201	1952	HKCSICHREFSTGHSLGGHKRLH
Q9SIJ0/65-87	1953	RPCTECGKQFGSLKALFGHMRCH
Q9SIJ0/148-170	1954	FECDGCKKVFGSHQALGGHRATH
Q9SIJ0/211-233	1955	HRCNICSRVFSSGQALGGHMRCH
Q9SLD4/47-69	1956	FECKTCNKRFSSFQALGGHRASH
Q9SLD4/94-116	1957	HKCSICSQSFGTQALGGHMRRH
Q9ZU93/121-143	1958	FECPICKNPFTSEEEVSVHVES
Q9SFT3/177-200	1959	CACPQCGEVFPKLESLEHHQAVRH
Q9ZQE0/244-266	1960	YTCPKCNGVFNTSQKFAAHMSSH
Q42423/80-102	1961	YKCSVCDKTFSSYQALGGHKASH
Q42423/136-158	1962	HVCTICNKSFPSPGQALGGHKRCH
Q9ZWA6/146-168	1963	WKCEKCAKRYAVQSDWKAHSKTC
Q9ZWA6/173-194	1964	YRCDCGTIFSRRDSFITHRAFC
Q9ZWA6/70-92	1965	FLCEICGKGQFQRDQNLQLHRRGH
O80942/39-61	1966	YTCSFCRREFKSAQALGGHMNVH
Q39217/90-112	1967	HKCSICSQSFGTQALGGHMRRH
Q39217/43-65	1968	FECKTCNKRFSSFQALGGHRASH
Q39092/160-182	1969	FECETCÉKVFKSYQALGGHRASH
Q39092/209-231	1970	HECPICAKVFTSGQALGGHKRSH
Q39092/5-27	1971	HKCKLCWKSFANGRALGGHMRSH
O81793/138-160	1972	PVCHICGRGFGGSWKAVF GHMRAH
O64828/530-553	1973	LQCIPCGSHFGDKEQLLVHVQAVH
O64828/599-622	1974	FVCKFCGLKFNLLPDLGRHHQAEH
O64828/496-519	1975	FACAICLDSFVRRKLLIEHVEERH
O49591/251-278	1976	FMCLYCNELCRPFSSLEAVRKHMEAKSH
O49591/26-50	1977	LTCNACNMEFKDEEERNLHYKSDWH
O49591/90-114	1978	YTCAIACAKGYRSSSKAHEQHLQSRSH

There follow several examples of how to construct and select DNA-binding sub-domains from libraries of natural zinc fingers.

5

Example 4: Human Zinc Finger Module ‘Mini-Library’.

As a preliminary test of the efficacy of using natural zinc finger modules for constructing novel DNA-binding domains, a ‘mini-library’ of natural, human zinc finger modules is

10 generated. The mini-library comprises 8 zinc finger modules, which have the following nomenclature assigned to them in the human genome database: Zif268 finger 1, Zif268 finger 2, Sp1 finger 3, WT1 finger 1, O15391, O75626, ZN45 and Z165. Since there is more than one zinc finger module belonging to the zinc fingers proteins ZN45 and Z165, we have called the selected modules ZN45-(AAA) and Z165-(GCC) respectively,

according to their predicted binding site. We have also predicted the binding sites for the zinc fingers O15391 and O75626. The preferred binding sites for Zif268 finger 1, Zif268 finger 2, Sp1 finger 3 and WT1 finger 1 are already known. The amino acid sequences of each of the stated modules, and their predicted or previously determined binding 5 sequences are shown in Table 3.

Two 3-zinc finger peptide libraries are prepared, containing the 8 zinc finger modules stated. All novel 3-finger peptides contain a leader sequence, MAEERP (SEQ ID NO:16), at the start of the peptide and are tagged by the sequence 10 LRQKDGGGSYPYDVPDYA (SEQ ID NO:1989) at the C-terminus. This sequence provides: (in the absence of a further C-terminal finger) a suitable terminus to the final α -helix of the peptide -LRQKD- (SEQ ID NO:1987) as found in wild-type Zif268; a short, flexible linker sequence, GGGS (SEQ ID NO:2121); and an HA-tag (YPYDVPDYA 15 (SEQ ID NO:2122)), which is recognised by the HA-antibody. Adjacent zinc finger modules are fused using the linker peptide sequence TGEKP (SEQ ID NO:3). The peptide sequences described above are also displayed in Table 3.

In the first library (library 1), the 8 zinc finger modules are recombined in random order to create 3-finger peptides with all possible combinations of the 8 zinc finger modules. 20 Such a procedure results in a library diversity of 512 ($=8^3$), comprising peptides that are predicted to bind to any possible combination of the binding sites assigned in Table 3. Library 1 allows novel 3-finger domains to be selected as a unit, for specified 9 bp target sequences. Such 3-finger units may be used for the construction of poly-zinc finger peptides as described in Moore, M., Choo, Y. & Klug, A. (2001) *Proc. Natl. Acad. Sci. 25 USA* 98: 1432-1436; and WO 01/53480.

In the second library (library 2), the 8 zinc finger modules are randomly recombined to create 2-finger peptides which are all joined to the C-terminus of Zif268 finger 1. The invariant finger 1 acts as an anchor for the selection, both by providing extra affinity to 30 stabilise the selection, and by fixing the register of the protein DNA interaction (as discussed *supra*). Such a library has a diversity of 64 ($=8^2$), and allows novel 2-finger units to be selected for a given 6 bp target sequence. The resulting 2 finger units can be

recovered by PCR and used in the construction of poly-zinc finger peptides (based on strings of 2-finger units), as described in WO 01/53480.

These two libraries (encoding 3-finger peptides) are screened, as described below, for the 5 ability of their encoded proteins to bind three different 9 bp binding sequences: 5'-GCG-TGG-GCG-3'; 5'-GGA-TAA-GCG-3'; and 5'-GCC-GAG-TGG-3'.

As positive controls, the genes encoding the 3-finger peptides predicted to bind the above target sequences are specifically constructed and tested in a similar manner.

10

x	FINGER/UNIT	SEQ ID NO:	PEPTIDE SEQUENCE	SITE
1	ZIF268 F1	1979	YACPVESCDRRFSRSDELTRHIRIH	GCG
2	ZIF268 F2	1980	FQCRIJCMRNFSRSRSDHLSTHIRTH	TGG
3	Sp1 F3	1981	FSCPICEKRFMRSDHLTKHARRH	GGG
4	WT1 F1	1982	FMCAYPGCNKRYFKLSHLQMHSRKH	GAG
5	O15391	1983	FVCPFDVCNRKFAQSTNLKTHILTH	TAA ¹
6	O75626	1984	FKCQTCNKGFTQLAHLQKHYLVH	GGA ¹
7	ZN45-AAA	1985	YKCEECGKGFSQASNLLAHQRGH	AAA ¹
8	Z165-GCC	1986	YECNECGKSFAESSDLTRHRRIH	GCC ¹
9	leader	16	MAEERP	-
10	linker	3	TGEKP	-
11	G ₃ S-HA-tag	1989	LRQKDGGGSYPYDVPDYA*	-

¹Predicted binding site. *indicates a translation stop codon.

Table 3. Nomenclature, amino acid sequences and known or predicted binding sequences ("SITE") of zinc finger modules and other peptide units used in library construction.

15

a. **Human Zinc Finger Mini-Library Construction.**

Two libraries are prepared, according to the scheme shown in Figure 2. The N-terminal finger of the 3-finger construct is referred to as 'cassette A'. The central finger is encoded by cassette B, and the third (C-terminal) finger module is called cassette C.

20

Zinc Finger Cassettes

Polynucleotide sequences encoding the amino acid sequences of the 8 zinc finger modules shown in Table 3 are determined, taking into account *E. coli* codon preferences,

and the corresponding nucleotide sequences are synthesised as single stranded oligonucleotides, examples of which are shown in Table 4. Also shown are the sequences of exemplary linkers and an exemplary 3'-tag required for the assembly of 3-finger domains. Double stranded cassettes encoding the zinc finger modules and relevant 5 leader, linker, and terminator sequences are generated by PCR according to the procedure described below, using the appropriate oligonucleotide templates of Table 4, and primers of Table 5.

x	CODE	FINGER	SEQ ID NO	NUCLEOTIDE SEQUENCE
1	AS144	ZIF268 F1	1990	TATGCGTGCCCGGTGGAAAGCTGCGATCGTCGTTTAG CCGTAGCGATGAAGTGCACCGTCATATTCTGATTCTAT
2	AS145	ZIF268 F2	1991	TTTCAGTGCCGTATTCGATGCGTAACCTTAGCCGTAG CGATCATCTGAGCACCCATATTCTGACCCAT
3	AS148	Sp1 F3	1992	TTTAGCTGCCGATTTGCGAAAAACGTTTATGCGTAG CGATCATCTGACAAACATGCGCGTCGTAT
4	AS149	WT1 F1	1993	TTTATGTGCGCGTATCCGGGCTGCAACAAACGTTATTT TAAACTGAGCCATCTGCAGatgcATAGCCGTAAACAT
5	AS150	O15391	1994	TTTGTGTGCCCGTTGATGTGTGCAACCGTAAATTGCG GCAGAGCACCAACCTGAAAACCCATATTCTGACCCAT
6	AS151	O75626	1995	TTTAAATGCCAGACCTGCAACAAAGGTTTACCCAGCT GGCGCATCTGCAGAAACATTATCTGGTGCAT
7	AS152	ZN45- AAA	1996	TATAAATGCGAAGAATCGGGCAAAGGTTTAGCCAGGC GAGCAACCTGCTGGCGCATCAGCGTGGCCAT
8	AS153	Z165-GCC	1997	TATGAATGCAACGAATCGGGAAAAGCTTGCAGAAAG CAGCGATCTGACCCGTATCGTCGTATTCT
9		MAEERP leader	1998	ATGGCGGAAGAACGTCCG
10		TGEKP linker	1999	ACCGGGAAAAACCG
11		G ₃ S-HA- tag (tag)	2000	CATCTGCGCCAGAAGGACGGCGGGCAGCTATCCGTA TGATGTGCCGGATTATGCGTAA

10 **Table 4.** Nucleotide sequences encoding zinc finger modules and other peptide sequences used in the construction of 3-finger proteins.

x	CODE	NAME	SEQ ID NO	SEQUENCE
1	AS5	pETFwd1	2001	CGCTGACTTCCCGCGTTCC
2	AS86	SDRev	2002	ATGTATATCTCCTTCTTAAAGTT
3	AS93	ZnF1Fwd	2003	AACTTTAAGAAGGAGATATACATATGGCGGAAGAA CGTCCGTATGCGTGCCCCGGTGGAAAG
4	AS94	ZnF2Fwd	2004	AACTTTAAGAAGGAGATATACATATGGCGGAAGAA CGTCCGTTTCAGTGCGTATTGCGATG

5	AS95	ZnF3Fwd	2005	AACTTTAAGAAGGAGATATACATATGGCGGAAGAA CGTCCGTTAGCTGCCGATTGCG
6	AS96	ZnF4Fwd	2006	AACTTTAAGAAGGAGATATACATATGGCGGAAGAA CGTCCGTTATGTGCGCGTATCCGGG
7	AS97	ZnF5Fwd	2007	AACTTTAAGAAGGAGATATACATATGGCGGAAGAA CGTCCGTTATGTGCGCGTATCCGGG
8	AS98	ZnF6Fwd	2008	AACTTTAAGAAGGAGATATACATATGGCGGAAGAA CGTCCGTTAAATGCCAGACCTGCAAC
9	AS99	ZnF7Fwd	2009	AACTTTAAGAAGGAGATATACATATGGCGGAAGAA CGTCCGTATAAATGCGAAGAATGCGGC
10	AS100	ZnF8Fwd	2010	AACTTTAAGAAGGAGATATACATATGGCGGAAGAA CGTCCGTATGAATGCAACGAATGCGGC
11	AS101	1Link1Rev	2011	CGGTTTTCGCCGGTATGAATAACGAATATGACGGG
12	AS102	1Link2Rev	2012	CGGTTTTCGCCGGTATGGGTACGAATATGGGTGC
13	AS103	1Link3Rev	2013	CGGTTTTCGCCGGTATGACGACGCGCATGTTGG
14	AS104	1Link4Rev	2014	CGGTTTTCGCCGGTATGTTACGGCTATGCATCTG
15	AS105	1Link5Rev	2015	CGGTTTTCGCCGGTATGGGTACGAATATGGGTTTC
16	AS106	1Link6Rev	2016	CGGTTTTCGCCGGTATGCACCAAGATAATGTTCTGC
17	AS107	1Link7Rev	2017	CGGTTTTCGCCGGTATGCCACGCTGATGCGC
18	AS108	1Link8Rev	2018	CGGTTTTCGCCGGTATGAATAACGACGATGACGGG
19	AS109	1Link1Fwd	2019	CATACGGCGAAAAACCGTATGCGTGCCTGGTGA AAG
10	AS110	1Link2Fwd	2020	CATACGGCGAAAAACCGTTCACTGCCGTATTG CATG
11	AS111	1Link3Fwd	2021	CATACGGCGAAAAACCGTTAGCTGCCGATTG CG
12	AS112	1Link4Fwd	2022	CATACGGCGAAAAACCGTTATGTGCGCGTATCC GGG
13	AS113	1Link5Fwd	2023	CATACGGCGAAAAACCGTTGTGTGCCGTTGA TGTG
14	AS114	1Link6Fwd	2024	CATACGGCGAAAAACCGTTAAATGCCAGACCTG CAAC
15	AS115	1Link7Fwd	2025	CATACGGCGAAAAACCGTATAAATGCGAAGAATG CGGC
16	AS116	1Link8Fwd	2026	CATACGGCGAAAAACCGTATGAATGCAACGAATG CGGC
17	AS117	2Link1Rev	2027	TGGCTTCTCACCCGTGTGATGAATACGAATATGAC GGGTC
18	AS118	2Link2Rev	2028	TGGCTTCTCACCCGTGTGATGGGTACGAATATGGG TGC
19	AS119	2Link3Rev	2029	TGGCTTCTCACCCGTGTGATGACGACGCGCATGTT TGG
20	AS120	2Link4Rev	2030	TGGCTTCTCACCCGTGTGATGTTACGGCTATGCA TCTG

21	AS121	2Link5Rev	2031	TGGCTTCTCACCCGTGTGATGGGTAGAAATATGGG TTTTC
22	AS122	2Link6Rev	2032	TGGCTTCTCACCCGTGTGATGCACCAAGATAATGTT TCTGC
23	AS123	2Link7Rev	2033	TGGCTTCTCACCCGTGTGATGCCACGCTGATGCG C
24	AS124	2Link8Rev	2034	TGGCTTCTCACCCGTGTGATGAATAACGACGATGAC GGG
25	AS125	2Link1Fwd	2035	CACGGGTGAGAAGCCATATGCGTCCCCGGTGGAAA G
26	AS126	2Link2Fwd	2036	CACGGGTGAGAAGCCATTCACTGCCGTATTCGA TG
27	AS127	2Link3Fwd	2037	CACGGGTGAGAAGCCATTAGCTGCCGATTTGCG
28	AS128	2Link4Fwd	2038	CACGGGTGAGAAGCCATTATGTGCGCGTATCCGG G
29	AS129	2Link5Fwd	2039	CACGGGTGAGAAGCCATTGTGTGCCGTTGATG TG
30	AS130	2Link6Fwd	2040	CACGGGTGAGAAGCCATTAAATGCCAGACCTGCA AC
31	AS131	2Link7Fwd	2041	CACGGGTGAGAAGCCATATAAATGCGAAGAATGCG GC
32	AS132	2Link8Fwd	2042	CACGGGTGAGAAGCCATATGAATGCAACGAATGCG GC
33	AS133	3HA1Rev	2043	CTAGGAATTCTTACGCATAATCCGGCACATCATA GGATAGCTGCCGCCGCCGTCTTCTGGCGCAGATG AATACGAATATGACGGGTC
34	AS134	3HA2Rev	2044	CTAGGAATTCTTACGCATAATCCGGCACATCATA GGATAGCTGCCGCCGCCGTCTTCTGGCGCAGATG GGTACGAATATGGGTGC
35	AS135	3HA3Rev	2045	CTAGGAATTCTTACGCATAATCCGGCACATCATA GGATAGCTGCCGCCGCCGTCTTCTGGCGCAGATG ACGACGCGCATTTGG
36	AS136	3HA4Rev	2046	CTAGGAATTCTTACGCATAATCCGGCACATCATA GGATAGCTGCCGCCGCCGTCTTCTGGCGCAGATG TTTACGGCTATGCATCTG
37	AS137	3HA5Rev	2047	CTAGGAATTCTTACGCATAATCCGGCACATCATA GGATAGCTGCCGCCGCCGTCTTCTGGCGCAGATG GGTCAGAATATGGTTTC
38	AS138	3HA6Rev	2048	CTAGGAATTCTTACGCATAATCCGGCACATCATA GGATAGCTGCCGCCGCCGTCTTCTGGCGCAGATG CACCAGATAATGTTCTGC
39	AS139	3HA7Rev	2049	CTAGGAATTCTTACGCATAATCCGGCACATCATA GGATAGCTGCCGCCGCCGTCTTCTGGCGCAGATG GCCACGCTGATGCGC
40	AS140	3HA8Rev	2050	CTAGGAATTCTTACGCATAATCCGGCACATCATA GGATAGCTGCCGCCGCCGTCTTCTGGCGCAGATG AATACGACGATGACGGG

41	AS141	Rev3	2051	CTAGGAATTCTTACGCATAATC
42	AS142	1LinkRev	2052	CGGTTTTCGCCGGTATG
43	AS143	2LinkRev	2053	TGGCTTCTCACCCGTGTG

Table 5. Modifying oligonucleotides used for mini-library construction.

5 1. *Library 1.*

Once made into double stranded DNA cassettes, the finger units are attached to T7 upstream expression sequences by PCR overlap extension, using the following protocol.

10 (a) Upstream sequences are first extracted from pET23a by PCR using primers pETFwd1 and SDRev, generating the fragment pET5'.

15 (b) The fingers for cassette A are amplified with forward primers ZnFxFwd (AS93-100) and reverse primers 1LinkxRev (AS101-AS108), where x is the number of a particular finger from Tables 3 and 4, as indicated.

20 (c) The fingers for cassette B are amplified with forward primers 1LinkxFwd (AS109-116) and reverse primers 2LinkxRev (AS117-AS124), where x refers to the finger module number.

25 (d) The fingers for cassette C are amplified with forward primers 2LinkxFwd (AS125-132) and reverse primers 3HAxRev (AS133-AS140), where x refers to the appropriate zinc finger module.

30 The steps to create cassettes A, B and C are performed separately, however, mixed populations of template oligonucleotides can be added to each PCR of steps (a), (b), and (c) to produce a library of each cassette.

The final 3-finger library is assembled by overlap extension as outlined in Figure 2. In 30 the first step the mixed pool of cassette A is appended to the upstream sequences, pET5'.

Equimolar amounts are mixed and PCR-cycled in the absence of primers. The reaction product is either purified immediately or reamplified before purification using primers pETFwd1 and 1LinkRev.

- 5 In the second step cassette B (mixed pool) is appended to the product of the above step. Again, equimolar amounts are mixed and PCR-cycled in the absence of primers. The reaction product is either purified immediately or reamplified before purification using primers pETFwd1 and 2LinkRev.
- 10 In the final step cassette C (mixed pool) is appended to the above product. Equimolar amounts are mixed and PCR-cycled in the absence of primers. As before, the reaction product may be purified immediately or reamplified before purification using primers pETFwd1 and Rev3. (see, also Figure 2).

15 2. *Library 2.*

Library 2 is assembled in a similar manner to Library 1 except that cassette A is represented by Zif268 finger 1 only.

- 20 The final PCR products containing T7 promoter sequences and encoding 3-finger peptides attached to an HA-antibody tag are purified and used for the production of protein.

25 b. **Zinc Finger Library Screening.**

Two exemplary methods for screening zinc finger libraries, such as those produced above, are described in Protocol A and Protocol B, below.

Protocol A:

The peptides of library 1 and library 2 are screened to select 3-zinc finger domains which bind the sequences: 5'-GCG-TGG-GCG-3'; 5'-GGA-TAA-GCG-3'; and 5'-GCC-GAG-5 TGG-3'. Since library 2 contains Zif268 finger 1 in the N-terminal position, in theory, these peptides should only bind the sequences, 5'-GCG-TGG-GCG-3', and 5'-GGA-TAA-GCG-3'. Hence, library 2 is effectively used to select 2-finger units which bind strongest to the 6 bp sequences, 5'-GCG-TGG-3', and 5'-GGA-TAA-3'. Double stranded binding sites for use in the selection protocol are generated by annealing the 10 complimentary oligonucleotides: Zif.b site and Zif site RC (AS154 and AS155); #1#5#6.b and #1#5#6 RC (AS156 and AS157); and #2#4#8.b and #2#4#8 RC (AS158 and AS159). The top strand of each binding site is biotinylated, allowing capture of binding site/zinc finger/HA-antibody ternary complexes to the streptavidin-coated plate in an ELISA screening assay. The oligonucleotides are displayed in Table 6, below.

15

x	Code	Name	SEQ ID NO	Sequence
1	AS154	Zif.b site	2054	TTTTTTTTTGGCGTGGCGTTTTTTTT
2	AS155	Zif site RC	2055	AAAAAAAAACGCCACGCAAAAAAAAAA
3	AS156	#1#5#6.b	2056	TTTTTTTTGGATAAGCGTTTTTTTT
4	AS157	#1#5#6 RC	2057	AAAAAAAAACGCTTATCCAAAAAAAAAA
5	AS158	#2#4#8.b	2058	TTTTTTTTGCCTGTTGGTTTTTTTT
6	AS159	#2#4#8 RC	2059	AAAAAAAAACCAACAGGCAAAAAAAAAA

Table 6. Oligonucleotide sequences used to generate double stranded binding sites used in the selection procedure.

20

The PCR-amplified 3-finger constructs are gel-purified from a 1% TAE-agarose gel using the Gel Extraction Kit (Qiagen) and quantified based on absorbance at 260 nM. Dilutions (in 0.25 mg/ml λ DNA) of DNA template encoding for either library 1 or 2 are prepared 25 at the final total template concentration of 4.2 fM and 1 fM, respectively. At these concentrations 1 μ l of template contains approximately 2500 and 600 molecules of library 1 or library 2, respectively. At such low concentrations, such samples must be PCR amplified to generate enough template for protein expression. Hence, these 1 μ l aliquots

are taken and added to 1 ml PCR pre-mix, containing primers Rev3 (AS141) and pETFwd2 (primer sequences shown below, see Table 7). The PCR pre-mixes are then aliquoted into 96 (or 384) well plates at 10 μ l per well, which is the equivalent of approximately 25 or 6 molecules of library 1 or library 2 template, respectively.

5 Templates are amplified using 30 cycles of PCR. After this first round of PCR, 0.5 μ l aliquots of PCR product are added to new 10 μ l PCR pre-mixes (in 96 or 384 well format), containing nested primers, pETFwd3 and Rev3, and amplified for another 30 cycles. The resultant product is concentrated enough to perform *in vitro* transcription / translation.

10 *In vitro* translation experiments using TNT PCR coupled transcription-translation mix (Promega) are assembled according to the manufacturer's instructions. Typically 5 μ l final volume contains 1 μ l of each PCR product and 4 μ l rabbit reticulocyte pre-mix (containing 20 μ M methionine, 12.5 μ g/ml λ *Hind* III digest (Roche), 500 μ M ZnCl₂ (Sigma), 0.7 μ l H₂O, 40 nM PCR-amplified DNA template). Reactions are incubated at 30°C for 90 minutes. 50 μ l PBS binding buffer containing 0.1 % BSA (Sigma), 0.5% Tween 20 (Sigma), 50 μ M ZnCl₂, 10 nM of the appropriate biotinylated binding site, 25 μ U/ml rat 3F10 anti-HA HRP conjugate (Roche) is added to the translation mix and incubated for 45 minutes at room temperature. The binding mix is thereafter transferred 20 to pre-blocked black streptavidin-coated 8-well strips or 96 / 384 well plates (Roche), and the ternary complexes containing 3-finger peptide, biotinylated binding site and anti-HA HRP antibody are captured while shaking at 200 rpm for 45 minutes at room temperature. The wells are then washed five times with 100 μ l PBS binding buffer containing 0.1 % BSA (Sigma), 0.5% Tween 20 (Sigma), 50 μ M ZnCl₂ to remove unbound components. 25 Finally, the retained HRP activity is measured by adding 50 μ l QuantaBlu fluorogenic HRP substrate (Pierce). Figure 3 demonstrates the capture and detection of target site-binding zinc finger peptides using the assay described. Fluorescence is measured on a SpectraMax Gemini XS (Molecular Devices) fluorescence microplate reader at 320 nm excitation, 433 nm emission and 420 nm cut-off values.

30 The wells that give the highest levels of fluorescence are those which contain the highest number of, or tightest binding 3-finger peptides. PCR products from the second PCR

amplification stage, corresponding to such samples, are purified from TAE-agarose gels and quantified, as above. Pure PCR products are diluted to approximately 50 molecules per μ l (which is equivalent to approximately 100 aM concentration) in 0.25 mg/ml λ DNA. As above, 1 μ l samples of template are added to 1 ml PCR pre-mix containing 5 primers, pETFwd4 and Rev3. 10 μ l aliquots are placed in each well of a 96 well plate. At this stage, there is (on average) 0.5 template molecules per aliquot. Therefore, generally speaking, half of the samples will contain no template and half will contain a single template molecule. Samples are then PCR amplified using 30 cycles. Again, 0.5 μ l PCR samples are taken from each well and amplified again by 30 cycles of PCR using 10 the nested primers, pETFwd5 and Rev3. 1 μ l of each of these PCR products is used for protein expression, as described above. At this stage, the highest levels of fluorescence correspond to the samples containing the tightest binding 3-finger peptides. The PCR product encoding such peptides is purified, as before, and can be sequenced to determine the protein sequence of the optimal 3-zinc finger domain for the appropriate binding site.

15 If further rounds of selection are required, PCR amplification can be conducted with the nested primers pETFwd6, pETFwd9 and pETFwd7, also shown below (Table 7).

NAME	SEQ ID NO	SEQUENCE
pETFwd1	2060	CGCTGACTTCCCGCGTTCC
pETFwd2	2061	TCCAGACTTACGAAACACGG
pETFwd3	2062	CGAAGACCATTACATGTTGTTGC
pETFwd4	2063	GTCGCAGACGTTGCAGC
pETFwd5	2064	GCAGTCGCTTCACGTTCGC
pETFwd6	2065	CGCTCGCGTATCGGTGATT
pETFwd9	2066	CATTCTGCTAACCAAGTAAGGC
pETFwd7	2067	GCCTAGCCGGGTCTCAAC

20 **Table 7:** Primers used for PCR amplification of 3-finger cassettes (as constructed by the procedure of Figure 2) to provide template used in screening zinc finger libraries.

Protocol B:

The peptides of library 2 were screened to select 3-zinc finger domains which bind the sequences: 5'-GCG-TGG-GCG-3', and 5'-GGG-AGG-CCT-3'. Double stranded binding sites for use in the selection protocol were generated by annealing the complementary oligonucleotides: Zif.b site and Zif site RC (AS154 and AS155, shown above), which generated the 5'-GCG-TGG-GCG-3' binding site; and the oligonucleotides 5'-
5 TTTTTTTTGGGAGGCCTTTTTTT-3' (SEQ ID NO:2123) and 5'-
10 AAAAAAAAAGGCCTCCAAAAAAA-3' (SEQ ID NO:2124), which generated the 5'-GGG-AGG-CCT-3' binding site. The top strand of each binding site was biotinylated, allowing capture of binding site/zinc finger/HA-antibody ternary complexes onto streptavidin-coated plate in an ELISA screening assay.

15 The 3-finger library 2 constructs were cloned into the multiple cloning site of vector pET23a (Novagen), using appropriate restriction sites. This library was then transformed into *E.coli* and plated out to grow single colonies. 384 colonies (which should represent the vast majority of the 64 member library) were picked into 2xYT media with ampicillin and cultures grown at 37°C overnight. Library 2 expression cassettes were recovered
20 from bacteria by PCR using primers pETFwdx (where x is 1-7, eg pETFwd1) and Rev3 as described in Protocol A above.

In vitro coupled transcription / translation of PCR products was conducted as described above, with the difference that each of the 384 zinc finger peptides was screened
25 individually in a well of a 384 well plate. The library was screened against the 5'-GCG-TGG-GCG-3', and 5'-GGG-AGG-CCT-3' binding sites, as detailed in Protocol A. Wells that yielded the highest levels of fluorescence were those which contain the tightest binding 3-finger peptides. The ELISA results from the screen of the 384 samples against the 5'-GCG-TGG-GCG-3' site are shown in Figure 4. Six constructs displayed
30 significant binding to the target site and these are termed C8, G16, I19, I23, J19 and K19 according to their coordinates on the 384-well plate. Similarly, one construct (B10)

showed strong binding to the 5'-GGG-AGG-CCT-3' target site. PCR products encoding the tightest binding peptides can be purified, as described *supra*, and sequenced.

5 Some of the selected constructs: C8, J19, K19, I23, G16 (which bind the 5'-GCG-TGG-GCG-3' site) and B10 (which binds the 5'-GGG-AGG-CCT-3' site), were selected and screened against a range of different binding sites to test their specificity. The sites used were: 5'-GCG-TGG-GCG-3'; 5'-CCA-CTC-GGC-3'; 5'-CCT-AGG-GGG-3'; 5'-GGA-TAA-GCG-3'; 5'-GGG-AGG-CCT-3'; 5'-GCG-TAA-GGA-3'; and 5'-GCG-GGG-GGA-3'. The binding assay was conducted as described above. The results (Figure 5) show that the selected 3-zinc finger peptides bind preferentially to their target site, in comparison to the alternative binding sites tested.

10

Example 5: Human Zinc Finger Module Libraries for Rapid Selection of 2-Finger Units.

15

The preferred subunits of a poly-zinc finger construction strategy are in the form of two-finger sub-domains. Assuming that there are 1,000 individual natural finger modules, a library of all combinations of such zinc finger modules, in 2-finger units, would contain 20 1,000,000 members. All of the 1,000 natural finger modules would have to be made from oligonucleotides, and the expense would be considerable. Furthermore, this figure is likely to be an underestimate of the number of natural fingers. Hence, due to the huge numbers of natural, human zinc finger modules available, it is advantageous to limit the size of the libraries screened, as discussed in the Description. One way in which library 25 size can be reduced is to limit the library members to zinc finger modules which are predicted to bind the desired sequence. For instance, based on the target sites in Example 1, if 2-finger domains are required to bind the sequence 5'-GCG-TGG-3', an individual library can be constructed from the zinc finger modules predicted to bind the sequences 5'-GCG-3' and 5'-TGG-3'. Equally, if the sequence 5'-GGA-TAA-3' is to be targeted, 30 zinc finger modules predicted to bind the sequences and 5'-GGA-3' and 5'-TAA-3' can be used. Table 8 shows the natural, human zinc finger modules from Example 1, which are predicted to bind the aforementioned 3 bp sequences.

5'-GCG-3'	5'-TGG-3'	5'-GGA-3'	5'-TAA-3'
Zif268 finger 1 (GCG)	Zif268 finger 2 (TGG)	BCL6 (NGA)	TYY1 (NAA)
Zif268 finger 3 (GCG)	MAZ finger 2 (TGG)	O75626 (GGA)	O15391 (YAA)
Sp1 finger 2 (GCG)	WT1 finger 3 (TGG)	ZN45 (N ^N /TA)	O75626 (YAA)
WT1 finger 4 (GCG)	SP4 (NGG)	O15535 (GNA)	ZN45 (N ^N /TA)
BTE1 (GCG)	BTE1 (NGG)	Q15776 (GNA)	Z136 (TNN)
O43296 (GNG)	Z136 (TNN)	O60893 (GNA)	Z239 (YAA)
Z174 (GCG, RNA)	Q15776 (NGG)	Z132 (a) (GGA)	Q15776 (a) (TNA)
Z202 (GCG, RNA)	ZN84 (YGG)	Z132 (b) (GGA)	Q15776 (b) (TNA)
		Z132 (GGN)	Z195 (YAA)
		ZN85 (GGA)	ZN84 (YAA)
			O75346 (TAA)
			ZN43 (TAA)

Table 8. The natural, human zinc finger modules predicted to bind the sequences 5'-GCG-3', 5'-TGG-3', 5'-GGA-3' and 5'-TAA-3'.

5

On the basis of the specificities shown in Table 5, a library of 2-finger units to target the 6 bp sequence 5'-GCG-TGG-3' has 64 (8x8) members, and a library to target the sequence 5'-GGA-TAA-3' has 120 (10x12) members. To screen sample sizes of this magnitude 10 we can construct each 2-finger unit specifically (using for example, an 8x8 or 10x12 matrix arrangement), and assay the samples containing individual clones using the fluorescent-ELISA protocol of Example 4. Such a procedure can save time in comparison to constructing all possible 64 or 120 variants in a random fashion (as a library), as described in Example 4, because the number of constructs screened would 15 have to be considerably higher.

a. Construction of 2-Finger Domains to Bind 5'-GCG-TGG-3'

A 64 member, 2-finger library is constructed from the natural, human zinc finger modules 20 predicted to bind the sequences 5'-GCG-3' and 5'-TGG-3' (Table 8, columns 1 and 2).

The 2-finger library units are all attached to the C-terminus of Zif268 finger 1, which acts as an anchor finger. The construction protocol is different from that described in Example 4, as described below.

5 **Zinc Finger Cassettes**

Nucleotide sequences encoding the amino acid sequences of the 16 zinc finger modules (Table 8, columns 1 and 2) are determined, taking into account human codon preferences, and the corresponding nucleotide sequences are synthesised as single stranded 10 oligonucleotides, shown in Table 9. Double stranded cassettes encoding the zinc finger modules and flanking linker sequences are generated by PCR using the appropriate primers, shown in Table 10.

X	FINGER	SEQ ID NO	NUCLEOTIDE SEQUENCE
1	Zif268 F1	2068	TACGCCTGCCCGTGGAGAGCTGCGACCGCCGTTCA CCGCAGCGACGAGCTGACCCGCCACATCCGATCCAC
2	Zif268 F3	2069	TTCGCCTGCGACATCTGCGGCCGCAAGTTCGCCCCGAG CGACGAGCGCAAGCGCCACACCAAGATCCAC
3	Sp1 F2	2070	TTCGCCTGCAGCTGGCAGGACTGCAACAAGAAGTTCGC CCGCAGCGACGAGCTGGCCGCCACTACCGCACCCAC
4	WT1 F4	2071	TTCAGCTGCCGCTGGCCCAGCTGCCAGAAGAAGTTCGC CCGCAGCGACGAGCTGGTGCGCCACCACAAACATGCAC
5	BTE1	2072	TTCCCCTGCACCTGGCCCGACTGCCTGAAGAAGTTCA CCGCAGCGACGAGCTGACCCGCCACTACCGCACCCAC
6	O43296	2073	TACGAGTGCCTGGAGTGCAGCAAGGCCTTCACCCGCAT GAGCGGCCTGACCCGCCACAAGCGCATCCAC
7	Z174	2074	TACAAGTGCAGCAGACTGCGGAAGAGCTTCACCTGGAA CAGCGAGCTGAAGCGCCACAAGCGCGTGCAC
8	Z202	2075	TACCGCTGCAGCAGACTGCGGAAGCACTTCCGCTGGAC CAGCGACCTGGTGCGCCACCAGCGCACCCAC
9	Zif268 F2	2076	TTCCAGTGCCGCATCTGCATGCCAACTTCAGCCGCAG CGACCACTGAGCACCCACATCCGACCCAC
10	MAZ F2	2077	TACAAGTGCAGCCACTGCGGAAGAGCTTCAGCCGCC CGACCACTGAACAGCCACGTGCAGGCCAGGTGCAC
11	WT1 F3	2078	TTCCAGTGCAAGACCTGCCAGCGCAAGTTCAGCCGCAG CGACCACTGAAGACCCACACCCGCACCCAC
12	Sp4	2079	CACAAGTGCCCTACAGCGGCTGCGGAAGGTGTACGG CAAGAGCAGCCACCTGAAGGCCACTACCGCGTGCAC
13	BTE1	2080	CACAAGTGCCCTACAGCGGCTGCGGAAGGTGTACGG CAAGAGCAGCCACCTGAAGGCCACTACCGCGTGCAC

14	Z136	2081	TTCGAGTGCAAGCGCTGCGCAAGGCCTTCCGAGCAG CAGCAGCTTCCGCTGCACGAGCGCACCCAC
15	Q15776	2082	TACGAGTGCGACGAGTGCGGCAAGACCTTCCGCCAG CAGCCACCTGATCGGCCACCAGCGCAGCCAC
16	ZN84	2083	TACGAGTGCGGCGAGTGCGGCAAGGCCTTCAGCCCAA GAGCCACCTGATCAGCCACTGGCGCACCCAC

¹ RNA Binding.

Table 9. Nucleotide sequences of zinc finger modules and nucleotide sequences encoding other peptide sequences used in the construction of peptides to bind the sequence 5'-GCG-TGG-3'.

The primers used to amplify the N-terminal finger of the pair (the equivalent of cassette B, above) add TGEKP (SEQ ID NO:3) linker sequences, and the restriction site *Xma*I (5'-CCC-GGG-3') at the 5' end and an *Age*I site (5'-ACC-GGT-3') at the 3' end. *Age*I and *Xma*I create compatible ends, but have unique restriction sites. These primers are called CasBxFwd and CasBxRev, respectively, where x refers to the number of the zinc finger module in Table 9. The primers used to amplify the C-terminal finger of the pair (the equivalent of cassette C, above) add TGEKP (SEQ ID NO:3) linker sequences, and the restriction site *Xma*I at the 5' end and a sequence encoding LRQKDGGGS (SEQ ID NO:2125), containing a restriction site for *Bam*HI at the 3' end. These primers are referred to as CasCxFwd and CasCxRev, respectively. The 16 individual zinc finger cassettes are then purified using the QIAquick PCR purification kit (Qiagen).

Name	SEQ ID NO	Sequence
CasB9Fwd	2084	GAT <u>CCCCGGGGAGAAGCCCTTCCAGTGCCGCATCTGCAT</u>
CasB10Fwd	2085	GAT <u>CCCCGGGGAGAAGCCCTACAACTGCAGCCACTGCGG</u>
CasB11Fwd	2086	GAT <u>CCCCGGGGAGAAGCCCTTCCAGTGCAAGACCTGCCA</u>
CasB12Fwd	2087	GAT <u>CCCCGGGGAGAAGCCCACAAGTGCCCTACAGCG</u>
CasB13Fwd	2088	GAT <u>CCCCGGGGAGAAGCCCACAAGTGCCCTACAGCG</u>
CasB14Fwd	2089	GAT <u>CCCCGGGGAGAAGCCCTCGAGTGCAAGCGCTGCG</u>
CasB15Fwd	2090	GAT <u>CCCCGGGGAGAAGCCCTACGAGTGCAGCGAGTGCG</u>
CasB16Fwd	2091	GAT <u>CCCCGGGGAGAAGCCCTACGAGTGCAGCGAGTGCG</u>
CasC1Fwd	2092	GAT <u>CCCCGGGGAGAAGCCCTACGCCCTGCCCGTGGAG</u>

CasC2Fwd	2093	GAT <u>CCCCGGGGAGAAGCCCTTCGCCCTGCGACATCTGCG</u>
CasC3Fwd	2094	GAT <u>CCCCGGGGAGAAGCCCTTCGCCCTGCGAGCTGGCAGG</u>
CasC4Fwd	2095	GAT <u>CCCCGGGGAGAAGCCCTTCAGCTGCCGCTGGCCC</u>
CasC5Fwd	2096	GAT <u>CCCCGGGGAGAAGCCCTTCCCCTGCACCTGGCCC</u>
CasC6Fwd	2097	GAT <u>CCCCGGGGAGAAGCCCTACGAGTGCCTGGAGTGCG</u>
CasC7Fwd	2098	GAT <u>CCCCGGGGAGAAGCCCTACAAGTGCACGACTGCGG</u>
CasC8Fwd	2099	GAT <u>CCCCGGGGAGAAGCCCTACCGCTGCACGACTGCG</u>
CasB9Rev	2100	CTTCTCAC <u>CGGTGTGGGTGCGGATGTGGGTG</u>
CasB10Rev	2101	CTTCTCAC <u>CGGTGTGCACCTGGCGCACGTG</u>
CasB11Rev	2102	CTTCTCAC <u>CGGTGTGGGTGCGGGTGTGGGT</u>
CasB12Rev	2103	CTTCTCAC <u>CGGTGTGCACGCGGTAGTGGGC</u>
CasB13Rev	2104	CTTCTCAC <u>CGGTGTGCACGCGGTAGTGGGC</u>
CasB14Rev	2105	CTTCTCAC <u>CGGTGTGGGTGCGCTCGTGCAG</u>
CasB15Rev	2106	CTTCTCAC <u>CGGTGTGGCTGCGCTGGTGGCC</u>
CasB16Rev	2107	CTTCTCAC <u>CGGTGTGGGTGCGCCAGTGGCT</u>
CasC1Rev	2108	GAT <u>CGGATCCGCCGCCGTCCCTCTGGCGCAGGTGGATGC</u> GGATGTGGCGG
CasC2Rev	2109	GAT <u>CGGATCCGCCGCCGTCCCTCTGGCGCAGGTGGATCT</u> TGGTGTGGCGC
CasC3Rev	2110	GAT <u>CGGATCCGCCGCCGTCCCTCTGGCGCAGGTGGGTGC</u> GGTAGTGGCG
CasC4Rev	2111	GAT <u>CGGATCCGCCGCCGTCCCTCTGGCGCAGGTGCATGT</u> TGTGGTGGCGC
CasC5Rev	2112	GAT <u>CGGATCCGCCGCCGTCCCTCTGGCGCAGGTGGGTGC</u> GGTAGTGGCG
CasC6Rev	2113	GAT <u>CGGATCCGCCGCCGTCCCTCTGGCGCAGGTGGATGC</u> GCTTGTGGCGG
CasC7Rev	2114	GAT <u>CGGATCCGCCGCCGTCCCTCTGGCGCAGGTGCACGC</u> GCTTGTGGCG
CasC8Rev	2115	GAT <u>CGGATCCGCCGCCGTCCCTCTGGCGCAGGTGGGTGC</u> GCTGGTGGCG

ScaRev	2116	GTCATGCCATCCGTAAGATGC
GSFwd	2117	GGCGGAT<u>CCCTATCCGTATGATGTG</u>
Zif1Fwd	2118	AGAGAGAGAGAG<u>ATCTATGGCGGAAGAACGTCCGTATGC</u> GTGCCCGGTGGAAAG
Zif1Rev	2119	AGCC<u>GGATCCC</u>AAAC<u>ACCGGTATGA</u>ATACGAATATGACG GG
pETRev1	2120	AGTGTAGCGGTACGCTGC

Table 10. Oligonucleotides used for PCR construction of rapid zinc finger library.
Annealing sequences are shown in bold, restriction sites are underlined.

5 *3-Finger Library Peptides*

The 2 natural zinc finger modules for each construct are appended to the C-terminus of Zif268 finger 1 (as in Example 4, library 2). Hence, a plasmid construct containing Zif268 finger 1 and appropriate restriction sites for cloning of the two natural finger 10 modules is also prepared. The construction and cloning procedure for the 3-finger libraries follows (see also Figure 6).

(a) The plasmid pET23a/TZF-HA was assembled by PCR amplification of plasmid pTFZ-KOX (described in co-owned WO 01/53480) with primers AS1 and AS2. 15 The sequences of these primers are as follows:

AS1: CGATGGATCCATGGGAGAGAAGGGCGCTGC (SEQ ID NO:2126)
AS2: GCGTAAAGCTTACGCATAATCCGGCACATCATACGGATAAGAG
CCGCCGCCGTCCTTCTGTCTAAATGGATT (SEQ ID NO:2127)

The PCR product was gel purified and digested with BamHI and HindIII, then 20 repurified and cloned into BamH I/Hind III-digested pET23a vector (Novagen), yielding pET23a/TFZ-HA. A number of clones were picked and sequenced to verify the correctness of the inserts.

(b) A fragment of approximately 1.2 kb is amplified from the vector 25 pET23a/TFZ-HA, using the primers ScaRev and GSFwd (Table 10). This fragment

contains the HA-epitope tag sequence (YPYDVPDYA* (SEQ ID NO: 2122)) and part of the GGGs (SEQ ID NO:1988) linker sequence at the 5' end. Additionally, the GSFwd primer adds a BamHI site at the extreme 5' end. The ScaRev primer does not contain a restriction site, but a *Scal* site from the vector is present approximately 40 bp downstream 5 of the primer binding site. This fragment is cut with *Bam*HI and *Scal* and inserted into similarly cut pET23a.

(c) Zif268 finger 1 is then amplified using the PCR primers Zif1Fwd and Zif1Rev (Table 10), which add a *Bgl*II site at the 5' end and both *Age*I and *Bam*HI sites at the 3' 10 end. This construct is then cut with *Bgl*II and *Bam*HI and inserted into the vector construct made in step (b), which has been linearised with *Bam*HI. At this stage the new construct, termed pET23aZif1HA is sequenced to find correctly oriented zinc finger inserts.

15 (d) Oligonucleotides encoding zinc finger modules for the C-terminus of the 3-finger constructs (cassette C) are amplified using the primers CasCxFor and CasCxRev (where x is 1 to 8, see Table 10). These cassettes are then digested with the restriction enzyme *Bam*HI, and inserted into *Bam*HI cut, dephosphorylated pET23aZif1HA. At this stage the new vector construct is not recircularised.

20 (e) Oligonucleotides encoding zinc finger modules for cassette B are amplified using primers CasBxFor and CasBxRev (where x is 9 to 16, see Table 10). These fragments are cut with the enzymes *Xma*I and *Age*I, at 37 °C for 1-2 hours. The linear vector produced in stage (d) above, is also cut with *Age*I and *Xma*I (as described), and 25 dephosphorylated. Digested cassette B fragments are ligated into *Age*I, *Xma*I cut vector, in the presence of the restriction enzymes *Age*I and *Xma*I at room temperature for 16 hours. During this incubation incorrectly ligated fragments are re-digested and re-ligated repeatedly, until the majority (or all) of the inserts are in the desired orientation. Correct 3-finger constructs have the assembly depicted in Figure 6.

30 (f) Finally, 3-finger constructs are amplified from the ligated vector (produced in step (e)) using the primers pETFwd1 (Table 5) and pETRev1 (Table 10). 1 µl of each

ligation mixture is amplified in a 10 μ l (total volume) PCR reaction for 30 cycles. Alternatively, the ligated vector can be transformed into bacteria to produce samples containing single zinc finger clones.

5 The above procedure results in the majority of PCR products being the correct 3-finger constructs, so that any incorrect fragments will not significantly affect the selection protocol, and the PCR products can be used for screening without further processing. Alternatively, 3-finger PCR products may be purified from an agarose gel before use.

10 **b. Screening of the Library Against 5'-GCG-TGG-GCG-3'**

Members of the zinc finger library can be screened against the desired target site from a mixed population of clones, or from individual clones as described in Example 4, Protocol A or Protocol B (above), respectively. The target site for the screen is produced 15 by annealing the oligonucleotides Zif.b site (AS154) and Zif site RC (AS155), as before. Template for protein expression is in each case made by PCR using primers pETFwd1 (Table 5) and pETRev1 (Table 10). 1 μ l of each PCR reaction is used to express protein and screen for binding to the Zif site in the manner described in Example 4. The DNA corresponding to the samples giving the highest fluorescence signals is collected, purified 20 from a 1% TAE-agarose gel, and sequenced to determine the sequence of the optimal binding 3-finger peptide.

Example 6: Reduced Human Zinc Finger Module Library for Universal DNA Recognition.

25 A library system similar to that described in Example 5 can be constructed using zinc finger modules from databases such as those in Examples 1, 2 and 3 to select 2-finger units which bind any 2-finger (6 bp) recognition sequence. There are only 4096 ($=4^6$) unique 6 bp sequences, therefore, a 2-finger library of natural zinc fingers (from specific 30 animals, plants or fungi) can easily be constructed with enough variability to provide a specific 2-finger combination for optimal binding to any 6 bp target site. Again, to reduce the number of natural zinc finger modules that have to be constructed, a small

selection of natural zinc finger modules (e.g., 3) are chosen for each 3 bp binding sequence (according to their predicted or determined recognition sequence). There are 64 ($=4^3$) possible 3 bp binding sequences so in the first instance less than 200 (i.e. 192) natural zinc finger modules are constructed. These 200 zinc finger modules can be in 5 either of 2 possible positions in the 2-finger construct, which gives approximately 40,000 ($=200^2$) combinations of fingers to bind the 4096 possible 6 bp target sites. As in Example 5, these 2-finger units are attached to Zif268 finger 1 which acts as an anchor for DNA recognition.

10 a. **Library Construction**

The selected zinc finger modules are reverse translated from their amino acid sequences and synthesised as oligonucleotides. Double stranded zinc finger cassettes for both N-terminal and C-terminal fingers are created by PCR using primers specific for the relevant 15 zinc finger module. Each zinc finger module is amplified in 2 separate reactions, as described in Example 5. The first PCR reaction uses primers which add TGEKP (SEQ ID NO:3) linker peptides and *Age*I and *Xma*I restriction sites, to the 3' and 5' ends, respectively, to generate cassette B fragments. The second PCR reaction generates cassette C fragments by adding a TGEKP (SEQ ID NO:3) linker and an *Xma*I site at the 20 5' end (this primer is the same as that used in cassette B production), and a sequence encoding the sequence LRQKDGGGS (SEQ ID NO:2125) and a *Bam*HI restriction site at the 3' end. The final constructs are similar to that represented in Figure 6.

25 b. **Library Selection**

The collection of 3-finger zinc finger peptides produced above can be used to obtain specific domains for binding desired target sequences. Two exemplary approaches are described below.

30 i). *Non-Cloning Selections.*

A library constructed as described herein can be used to select optimal zinc finger domains for binding to any specified binding site. For instance, to select a peptide which binds the sequence 5'-GGA-TAA-3', the binding site formed by annealing the 5 oligonucleotides #1#5#6.b and #1#5#6 RC (Table 6, above), can be used as a target site (5'-GGA-TAA-GCG-3'). Selection of a zinc finger domain to bind such a target can be conducted, for example, in the manner described in Example 4. Briefly, the zinc finger library is diluted into 100 or more sub-libraries, which are screened as described above. The most active sub-libraries collected are further diluted to create much smaller sub-libraries, which are screened again, and so on. Following such a protocol, a library of 10 40,000 members can be fully screened and a high-affinity binder selected in just 3 rounds.

This selection procedure provides an extremely rapid method to select zinc finger peptides to bind any desired target site. The procedure also has the advantages of 15 eliminating the need for cloning (as is required for methods such as phage display, see below), and is not limited by library size.

ii). Phage Library Selections

Zinc finger polypeptide phage display libraries are made and used to select clones 20 encoding peptides that bind the desired nucleotide sequence, as described in co-owned WO 98/53057. An exemplary phage display library contains peptides which bind target sites with the sequence 5'-XXX-XXX-GCG-3', where X can be any nucleotide. Hence, libraries of phage can be selected using the same target sites as described above. The selection protocol for zinc fingers displayed on phage is briefly described below.

25

Protocol

The selection protocol is adapted from that described in co-owned international patent application WO98/53057.

30

The 3-finger constructs of the present Example are PCR amplified using universal forward and reverse primers which contain sites for *NotI* and *SfiI* respectively (called NatPhageF and NatPhageR, respectively).

5 NatPhageF: GCAACTGCGGCCCAGCCGGCCATGGCAGAGGAACGCCCGTATG (SEQ ID
NO:2128)
NatPhageR: GAGTCATTCTGCGGCCGCTCCTTCTGGCGCAGGTG (SEQ ID NO:2129)

Backward PCR primers in addition introduce Met-Ala-Glu as the first three amino acid
10 residues of the zinc finger polypeptides, and these are followed by the residues of the
wild type or library zinc finger polypeptides as required. Cloning overhangs are
produced by digestion with *SfiI* and *NotI* where necessary. Nucleic acid encoding zinc
15 finger polypeptide fragments is ligated into similarly prepared Fd-Tet-SN vector. This is
a derivative of fd-tet-DOG1 (Hoogenboom *et al.* (1991) *Nucl. Acids Res.* 19:4133-4137),
in which a section of the pelB leader and a restriction site for the enzyme *SfiI* (underlined)
have been added by site-directed mutagenesis using the oligonucleotide:

5' CTCCTGCAGTTGGACCTGTGCCATGGCCGGCTGGGCCGCATA
GAATGGAAACAACTAAAGC 3' (SEQ ID NO:2130)

20 that anneals in the region of the polylinker. Electrocompetent DH5 α cells are
transformed with recombinant vector in 200 ng aliquots, grown for 1 hour in 2xTY
medium with 1% glucose, and plated on TYE containing 15 μ g/ml tetracycline and 1%
glucose.

25 To generate phage for selections, tetracycline resistant colonies are transferred from
plates into 2xTY medium (16g/litre Bacto tryptone, 10g/litre Bacto yeast extract, 5g/litre
NaCl) containing 50 μ M ZnCl₂ and 15 μ g/ml tetracycline, and cultured overnight at 30°C
in a shaking incubator. Cleared culture supernatant containing phage particles is obtained
by centrifuging at 300 xg for 5 minutes.

Double stranded binding sites for use in selections are generated by annealing complementary oligonucleotides, one of which is biotinylated.

Biotinylated DNA target sites (1 pmol) are bound to streptavidin-coated wells (Roche).

5 Phage supernatant solutions are diluted 1:10 in PBS selection buffer (PBS containing 50 μ M ZnCl₂, 2% Marvel, 1% Tween, 20 μ g/ml sonicated salmon sperm DNA, and 10-fold excess of competitor DNA), and 200 μ l is applied to each well for 1 hour at 20°C. After this time, the wells are emptied and washed 18 times with PBS containing 50 μ M ZnCl₂ and 1% Tween and 2 times in PBS containing 50 μ M ZnCl₂. Retained phage are eluted in

10 100 μ l 0.1M triethylamine and neutralised with an equal volume of 1M Tris (pH 7.4).

Logarithmic-phase *E. coli* JM109 (100 μ l) are infected with eluted phage (100 μ l), and used to prepare phage supernatants for subsequent rounds of selection. After 4 rounds of selection, a 'pool' or 'mini-population' of phage is obtained, which bind the specified target sequence. These pools of phage can be stored at -70°C for later use. Additionally,

15 15 *E. coli* infected with these phage pools can be plated to obtain individual clones, which can be tested by ELISA for binding affinity and specificity to obtain the 'best' clone (see Example 9, Quality Control).

20 **Example 7: Complete Human Zinc Finger Module Library for Universal DNA Recognition.**

An complete, or nearly complete, library containing all zinc finger sequences which bind a particular target site can be constructed using zinc finger modules to select 2-finger (or 25 3-finger) units which bind any 6 bp (or 9 bp) recognition sequence. Two exemplary methods for construction of such a library are described.

a. **Oligonucleotide-Based Library Construction.**

30 All zinc finger modules may be synthesised as a single stranded oligonucleotide, as described in Example 4. Zinc finger modules are made double stranded and TGEKP (SEQ ID NO:3) linkers added by PCR with 5' and 3' primers specific for each individual

zinc finger module, to make cassettes. These cassettes can then be recombined, as described in Example 5, to make random or deliberate combinations of zinc finger modules comprising 2, 3, or more linked fingers.

5 **b. PCR-Based Library Construction.**

Zinc fingers proteins (especially of the Cys₂His₂ family) form the second most abundant family of proteins in the human genome. Furthermore, in nature, zinc finger modules are often linked by the canonical linker peptide TGEKP (SEQ ID NO:3), which begins 10 immediately after the second zinc-coordinating histidine residue. Therefore, the peptide sequence HTGEKP (SEQ ID NO:2131) is commonly found between natural zinc finger modules. Because of this consensus sequence, it has been possible to clone natural zinc finger modules from the human genome (Becker, K.G., Nagel, J.W., Canning, R.D., Biddison, W.E., Ozato, K. & Drew, P.D. (1995) *Hum. Mol. Genet.* 4: 685-691; Bray, P., 15 Lichter, P., Thiesen, H.-J., Ward, D.C. & Dawid, I.B. (1991) *Proc. Natl. Acad. Sci. USA* 88: 9563-9567), and the *Arabidopsis* genome (Meissner, R. & Michael, A.J. (1997) *Plant Mol Biol* 33: 615-624), using redundant primers for PCR. *See also Pellegrino et al.* 20 (1991) *Proc. Natl. Acad. Sci. USA* 88:671-675. It is preferable to use genomic DNA or a genomic DNA (gDNA) library, rather than a cDNA library, because transcription factors, such as zinc finger proteins, are strongly regulated during the cell cycle, development and 25 in response to extracellular signals. Hence, a cDNA library will probably not contain the majority of zinc finger proteins, and will be biased towards highly expressed proteins.

A suitable protocol for the PCR-extraction of zinc finger modules from human genomic 25 DNA follows:

Genomic DNA is purified directly from human cells, or provided by a gDNA library. gDNA libraries are preferable as they are commercially available (for example from Clontech, ATCC, Stratagene etc) and can be easily manipulated. PCR to extract zinc 30 finger modules can be conducted directly on purified gDNA, or the gDNA library can be screened for zinc fingers containing the HTGEKP (SEQ ID NO:2131) motif before carrying out PCR. To screen the gDNA library, any method known to one of skill in the

art, *e.g.* colony hybridisation, can be used. Phage containing gDNA inserts are plated onto *Escherichia coli* XL-1 Blue bacterial lawns. At least 10^6 phage plaques are transferred to replica filters and screened with, for example, a 27-mer ^{32}P -radiolabelled degenerate oligonucleotide, which anneals to the conserved linker region of zinc finger proteins and adjacent sequences. The sequence of a suitable degenerate probe (SEQ ID NO:2132), and the amino acid sequence (SEQ ID NO:2133) to which it corresponds is shown below.

10 $\text{C}^{\text{G}}/\text{T}^{\text{C}}/\text{G}$ $\text{A}^{\text{T}}/\text{C}^{\text{C}}/\text{G}$ $\text{CA}^{\text{C}}/\text{T}$ $\text{AC}^{\text{C}}/\text{G}$ $\text{GG}^{\text{C}}/\text{G}$ $\text{GA}^{\text{G}}/\text{A}$ $\text{AA}^{\text{G}}/\text{A}$ $\text{CC}^{\text{C}}/\text{T}$ $\text{T}^{\text{A}}/\text{T}^{\text{C}}/\text{T}$
R/L I/T/M H T G E K P Y/F

15 Hybridisation is performed, *e.g.*, for 16 hours at 42-50 °C, following which filters are washed 3-5 times, to remove non-specifically bound probe, in 0.2x standard saline citrate (SSC)/0.1% SDS. Filters are then subjected to autoradiography or phosphorimaging to determine positive plaques.

Positive plaques are picked into log-phase *E. coli* XL-1 Blue bacterial cultures and the phage are harvested for PCR. 1 μl phage supernatant is added to 49 μl PCR pre-mix, containing the oligonucleotide primers TGEKPfor (SEQ ID NO:2134) and TGEKPrev (SEQ ID NO:2135) (shown below, annealing sequence in bold), and zinc finger modules are amplified by 30 cycles of PCR. TGEKPfor (SEQ ID NO:2134) and TGEKPrev (SEQ ID NO:2135) also contain *Xba*I and *Eco*RI restriction sites (underlined), respectively. PCR products are separated on 1.5% TAE-agarose gels and fragments of approximately 120 bp (corresponding to 1 zinc finger module plus flanking sequences) are purified, as described in Example 4. Additionally, fragments of approximately 220 bp, corresponding to natural 2-finger units, can also be collected and used. Such products can be digested with *Xba*I and *Eco*RI and cloned into a vector that has been digested so as to generate compatible ends, such as, for example, pcDNA3.1(-) (Invitrogen) digested with *Eco*RI and *Xba*I. Such a vector pool can then be used as a source for natural 1- or 2-zinc finger modules, from which to construct 2- or 3-zinc finger peptides for selections as described above. Zinc finger modules for cassette B can be amplified from such vectors using the universal primers TGEKPXma (SEQ ID NO:2136) and TGEKPAge (SEQ ID NO:2137),

which anneal to the conserved TGEKP (SEQ ID NO:3) linker regions and add restriction sites for the enzymes *Xma*I at the 5' terminus and *Age*I at the 3' terminus, respectively (restriction sites underlined). Cassette C units can be amplified using the primer TGEKPxma (SEQ ID NO:2136) and TGEKpend (SEQ ID NO:2138), which adds a 3' 5 TRQKDGGGS (SEQ ID NO:2139) sequence incorporating a *Bam*HI site (underlined, see below). Two- and 3-finger constructs can then be constructed and screened as described in the Examples above.

TGEKPfor: TTAGTCTAGA^C/_GCA^C/_TAC^C/_GGG^C/_GGA^G/_AAA^G/_ACC (SEQ ID 10 NO:2134)

TGEKPrev: TACTGAATTC^G/_AGG^C/_TTT^C/_TTC^G/_CCC^G/_CGT^G/_ATG (SEQ ID NO:2135)

TGEKPxma: TCTAGA^C/_GCA^C/_TCCCGGGG^G/_AAA^G/_ACC (SEQ ID NO:2136)

TGEKPAge: GAATTC^G/_AGG^C/_TTT^C/_TTCACCGGT^G/_ATG (SEQ ID NO:2137)

15 TGEKpend: AGTGTGGTGGAATTC^G/_AGGGGATCCGGCCGCCGTC^C/_TTT^C/_TTG^G/_CCG^G/_CGT^G/_ATG (SEQ ID NO:2138)

Example 8. Microarray Analysis.

20 Microarray analysis can also be used to determine the binding site specificity of 2- and 3-finger peptides. For example, a 3-zinc finger library, with finger 1 fixed as Zif268 finger one recognises the sequence 5'-XXX-XXX-GCG-3', where X is any specified nucleotide. Hence, there are 4096 ($=4^6$) unique binding sites for such a library. All 4096 of these 25 sites can be arrayed onto a single glass slide, allowing a specified 2-finger peptide to be screened against every possible binding site at once. A suitable protocol for such an experiment is described in Martha L. Bulyk, Xiaohua Huang, Yen Choo, & George M. Church (*Proc. Natl. Acad. Sci. USA*: Vol. 98, No. 13, 7158-7163, June 19, 2001) which is incorporated, by reference, in its entirety. See also co-owned WO 01/25417, the 30 disclosure of which is hereby incorporated by reference in its entirety.

The amount of binding to each target sequence can be visualised and quantified using simple fluorescence measurements. For example, the zinc finger peptide can be expressed *in vitro*, or on the surface of phage. Isolated zinc finger peptides may contain an epitope tag for labelling purposes, whereas bound phage can be detected using a 5 primary antibody against a phage coat protein, such as gVIII. A secondary antibody, such as one conjugated to R-phycoerythrin may be used to provide a visible signal when a suitable substrate is applied.

10 **Example 9. Quality Control.**

Particular 2- or 3-finger peptides can be screened to determine their specificity or affinity, as desired.

a. **Phage ELISA Assay**

Phage supernatants from Round 4 of selection (Example 6, *supra*) are used to infect *E. coli* JM109 bacteria, and grown to prepare fresh supernatants for zinc finger phage

5 ELISA, using standard procedures as described previously (Choo, Y. & Klug, A. (1994) *Proc. Natl. Acad. Sci. USA* 91, 11163-11167; Choo, Y. & Klug, A. (1994) *Proc. Natl. Acad. Sci. USA* 91, 11168-11172). Briefly, 5'-biotinylated, positionally randomised oligonucleotide libraries, containing Zif268 binding site variants, are synthesised by annealing complimentary oligonucleotides as described *supra*. DNA libraries are added
10 to streptavidin-coated ELISA wells (Boehringer-Mannheim) in PBS containing 50µM ZnCl₂ (PBS/Zn). Phage solution (overnight bacterial culture supernatant diluted 1:10 in PBS/Zn containing 2% Marvel, 1% Tween and 20µg/ml sonicated salmon sperm DNA) is applied to each well (50µl/well). Binding is allowed to proceed for one hour at 20°C. Unbound phage are removed by washing 7 times with PBS/Zn containing 1% Tween,
15 then 3 times with PBS/Zn. Bound phage are detected by ELISA using horseradish peroxidase-conjugated anti-M13 IgG (Pharmacia Biotech) and the colourimetric signal is quantitated using SOFTMAX 2.32 (Molecular Devices).

For rapid validation, the entire population of phage from Round 4 selection can be

20 assayed in two ELISA wells: one containing the target DNA binding site, and one containing a control DNA binding site with between 1 and 5 base changes from the target sequence. A selection is deemed to be successful if the ELISA signal (representing DNA binding) is higher in the target well than in the control well.

25 The higher the signal measured above, the greater the *population* of specific binding clones. However, individual low values for such a procedure do not necessarily indicate a failure of the selection, as there may be individual high affinity / specificity clones within the round 4 phage population that may be masked by other non-specific clones. Nevertheless, this assay provides a quick profile of the overall quality of selection.

For a more detailed validation, individual phage clones are recovered from Round 4 by plating out infected bacterial colonies on agar. Fresh phage supernatants are prepared from these colonies and assayed by ELISA, as described above.

5 Finally, the coding sequence of individual zinc finger clones can be amplified by PCR using external primers complementary to phage sequence, and the PCR products are then sequenced to determine the amino acid sequence of the selected zinc fingers.

10 As an alternative, individual 3-finger peptides can be analysed by gel-shift assays or by microarray screening, as described *infra*. See also WO 00/41566, WO 00/42219 and WO 01/25417.

b. Gel-Shift Assay

15 Peptides are assayed using ^{32}P end-labelled synthetic oligonucleotide duplexes containing the appropriate binding site sequences.

DNA binding reactions contain the appropriate zinc-finger peptide, binding site and 1 μg competitor DNA (*e.g.*, poly dI-dC or salmon sperm DNA) in a total volume of 10 μl , which contains: 20 mM Bis-tris propane (pH 7.0), 100 mM NaCl, 5 mM MgCl₂, 50 μM ZnCl₂, 5 mM DTT, 0.1 mg/ml BSA, 0.1% Nonidet P40. Incubations are performed at 20 room temperature for 1 hour.

To determine the concentration of zinc finger peptide produced in the *in vitro* expression system, crude protein samples are used in gel-shift assays against a dilution series of the appropriate binding site. Binding site concentration is always well above the Kd of the peptide, but ranged from a higher concentration than the peptide (80 mM), at which all 25 available peptide binds DNA, to a lower concentration (3-5 mM), at which all DNA is bound. Controls are carried out to ensure that binding sites are not shifted (*i.e.*, bound) in the absence of zinc finger peptide. The reaction mixtures are then separated on a 7% native polyacrylamide gel. Radioactive signals are quantitated by PhosphorImager

analysis to determine the amount of shifted binding site, and hence, the concentration of active zinc finger peptide.

Dissociation constants (K_d) are determined in parallel to the calculation of active peptide concentration. For determination of K_d , serial 3, 4 or 5-fold dilutions of crude peptide are 5 made and incubated with radiolabelled binding site (10 pM – 10 nM depending on the peptide), as above. Samples are run on 7% native polyacrylamide gels and the radioactive signals quantitated by PhosphorImager analysis. The data is then analysed according to linear transformation of the binding equation and plotted in CA-Cricket Graph III (Computer Associates Inc. NY) to generate the apparent dissociation constants.

10 The K_d values reported are the average of at least two separate determinations.

c. **Microarray Assay**

15 Selected zinc finger domains can also be assayed for binding site specificity using the microarray analysis outlined in Example 8.

All publications mentioned in the above specification are herein incorporated by reference. Various modifications and variations of the described methods and system of the invention will be apparent to those skilled in the art without departing from the scope 20 and spirit of the invention. Although the invention has been described in connection with specific preferred embodiments, it should be understood that the invention as claimed should not be unduly limited to such specific embodiments. Indeed, various modifications of the described modes for carrying out the invention which are apparent to those skilled in molecular biology or related fields are intended to be within the scope of 25 the following claims.

CLAIMS

1. A composite binding polypeptide comprising a first natural binding domain derived from a first natural binding polypeptide, and a second natural binding domain derived from a second natural binding polypeptide, wherein said first and second natural binding polypeptides may be the same or different; which polypeptide binds to a target, said target differing from the natural target of the both the first and the second binding polypeptides.
2. A composite polypeptide according to claim 1, wherein said first and second natural binding polypeptides are different polypeptides.
3. A composite polypeptide according to claim 1 or claim 2, comprising three or more natural binding domains.
4. A composite polypeptide according to any preceding claim, wherein the binding domains are nucleic acid binding domains.
5. A composite polypeptide according to claim 4, which is a nucleic acid binding polypeptide.
6. A composite polypeptide according to claim 4 or claim 5 which is a zinc finger polypeptide, and the natural binding domains are zinc finger domains.
7. A composite polypeptide according to claim 6, which comprises a Cys2-His2 zinc finger binding domain.
8. A composite polypeptide according to claim 6 or claim 7, which comprises a Cys3-His zinc finger binding domain.
9. A composite polypeptide according to any preceding claim, which comprises 6 or more natural binding domains.

10. A composite polypeptide according to claim 9, wherein 6 natural binding domains are arranged in a 3x2 conformation, separated by linker sequences.
11. A chimeric polypeptide comprising:
 - (a) a binding polypeptide according to any preceding claim, and
 - (b) a biological effector domain.
11. A library of natural binding domains.
12. A library according to claim 11, comprising a plurality of natural binding domains from which a polypeptide according to any one of claims 1 to 10 can be assembled.
13. A library of natural zinc finger nucleic acid binding domains, wherein said zinc finger domains comprise a linker attached thereto.
14. A library according to claim 13, wherein the linker comprises the sequence TGEKPK.
15. A method for selecting a binding polypeptide capable of binding to a target site, comprising:
 - (a) providing a library of natural binding domains;
 - (b) assembling two or more of said domains to form a composite polypeptide;
 - (c) screening said composite polypeptide against the target site in order to determine its ability to bind the target site.
16. A method according to claim 15, wherein the natural binding domains are zinc finger binding domains.
17. A method according to claim 15 or claim 16, wherein two or more composite polypeptides comprising two or more domains which are selected for binding to two or

more target sites are combined to provide a composite polypeptide which binds to an aggregate binding site comprising the two or more target binding sites.

18. A method for designing a composite binding polypeptide, comprising:

(a) providing information defining a target site;

(b) selecting, from a database of natural binding domains, sequences of binding domains which are predicted to bind to the target site by the application of one or more rules which define target binding interactions for the binding domains; and

(c) displaying the sequences of the binding domains, separated by linker sequences, and optionally assembling the binding polypeptide from a library of said domains.

19. A method according to claim 18, wherein the binding domains are zinc finger domains.

20. A method according to claim 19, wherein the zinc fingers are considered to bind to a nucleic acid triplet and domains are selected according to one or more of the following rules:

(a) if the 5' base in the triplet is G, then position +6 in the α -helix is Arg; or position +6 is Ser or Thr and position ++2 is Asp;

(b) if the 5' base in the triplet is A, then position +6 in the α -helix is Gln and ++2 is not Asp;

(c) if the 5' base in the triplet is T, then position +6 in the α -helix is Ser or Thr and position ++2 is Asp;

(d) if the 5' base in the triplet is C, then position +6 in the α -helix may be any amino acid, provided that position ++2 in the α -helix is not Asp;

(e) if the central base in the triplet is G, then position +3 in the α -helix is His;

(f) if the central base in the triplet is A, then position +3 in the α -helix is Asn;

(g) if the central base in the triplet is T, then position +3 in the α -helix is Ala, Ser or Val; provided that if it is Ala, then one of the residues at -1 or +6 is a small residue;

(h) if the central base in the triplet is C, then position +3 in the α -helix is Ser, Asp, Glu, Leu, Thr or Val;

(i) if the 3' base in the triplet is G, then position -1 in the α -helix is Arg;

(j) if the 3' base in the triplet is A, then position -1 in the α -helix is Gln;

(k) if the 3' base in the triplet is T, then position -1 in the α -helix is Asn or Gln;

(l) if the 3' base in the triplet is C, then position -1 in the α -helix is Asp.

21. A method according to claim 19, wherein the zinc fingers are considered to bind to a nucleic acid quadruplet and domains are selected according to one or more of the following rules:

(a) if base 4 in the quadruplet is G, then position +6 in the α -helix is Arg or Lys;

(b) if base 4 in the quadruplet is A, then position +6 in the α -helix is Glu, Asn or Val;

(c) if base 4 in the quadruplet is T, then position +6 in the α -helix is Ser, Thr, Val or Lys;

(d) if base 4 in the quadruplet is C, then position +6 in the α -helix is Ser, Thr, Val, Ala, Glu or Asn;

(e) if base 3 in the quadruplet is G, then position +3 in the α -helix is His;

(f) if base 3 in the quadruplet is A, then position +3 in the α -helix is Asn;

(g) if base 3 in the quadruplet is T, then position +3 in the α -helix is Ala, Ser or Val; provided that if it is Ala, then one of the residues at -1 or +6 is a small residue;

(h) if base 3 in the quadruplet is C, then position +3 in the α -helix is Ser, Asp, Glu, Leu, Thr or Val;

(i) if base 2 in the quadruplet is G, then position -1 in the α -helix is Arg;

(j) if base 2 in the quadruplet is A, then position -1 in the α -helix is Gln;

(k) if base 2 in the quadruplet is T, then position -1 in the α -helix is His or Thr;

(l) if base 2 in the quadruplet is C, then position -1 in the α -helix is Asp or His;

(m) if base 1 in the quadruplet is G, then position +2 is Glu;

(n) if base 1 in the quadruplet is A, then position +2 Arg or Gln;

(o) if base 1 in the quadruplet is C, then position +2 is Asn, Gln, Arg, His or Lys;

(p) if base 1 in the quadruplet is T, then position +2 is Ser or Thr.

22. A method according to claim 19, wherein the zinc fingers are considered to bind to a nucleic acid quadruplet and domains are selected according to one or more of the following rules:

- (a) if base 4 in the quadruplet is G, then position +6 in the α -helix is Arg; or position +6 is Ser or Thr and position ++2 is Asp;
- (b) if base 4 in the quadruplet is A, then position +6 in the α -helix is Gln and ++2 is not Asp;
- (c) if base 4 in the quadruplet is T, then position +6 in the α -helix is Ser or Thr and position ++2 is Asp;
- (d) if base 4 in the quadruplet is C, then position +6 in the α -helix may be any amino acid, provided that position ++2 in the α -helix is not Asp;
- (e) if base 3 in the quadruplet is G, then position +3 in the α -helix is His;
- (f) if base 3 in the quadruplet is A, then position +3 in the α -helix is Asn;
- (g) if base 3 in the quadruplet is T, then position +3 in the α -helix is Ala, Ser or Val; provided that if it is Ala, then one of the residues at -1 or +6 is a small residue;
- (h) if base 3 in the quadruplet is C, then position +3 in the α -helix is Ser, Asp, Glu, Leu, Thr or Val;
- (i) if base 2 in the quadruplet is G, then position -1 in the α -helix is Arg;
- (j) if base 2 in the quadruplet is A, then position -1 in the α -helix is Gln;
- (k) if base 2 in the quadruplet is T, then position -1 in the α -helix is Asn or Gln;
- (l) if base 2 in the quadruplet is C, then position -1 in the α -helix is Asp;
- (m) if base 1 in the quadruplet is G, then position +2 is Asp;
- (n) if base 1 in the quadruplet is A, then position +2 is not Asp;
- (o) if base 1 in the quadruplet is C, then position +2 is not Asp;
- (p) if base 1 in the quadruplet is T, then position +2 is Ser or Thr.

23. The method of any of claims 18-22, further comprising the step of synthesizing a polynucleotide encoding the binding polypeptide.

24. A computer-implemented method for designing a zinc finger polypeptide, comprising the steps of:

- (a) providing a system comprising at least storage means for storing data relating to a library of zinc fingers; storage means for storing a rule table; means for inputting target nucleic acid sequence data; processing means for generating a result; and means for outputting the result;
- (b) inputting sequence data for a target nucleic acid molecule;
- (c) defining a first target zinc finger binding site in said nucleic acid molecule;
- (d) interrogating the zinc finger library and rule table storage means, comparing zinc fingers to the target zinc finger binding site according to the rule table and selecting zinc finger data identifying a zinc finger capable of binding to said target site;
- (e) defining at least one further target zinc finger binding site and repeating step (d); and
- (f) outputting the selected zinc finger data.

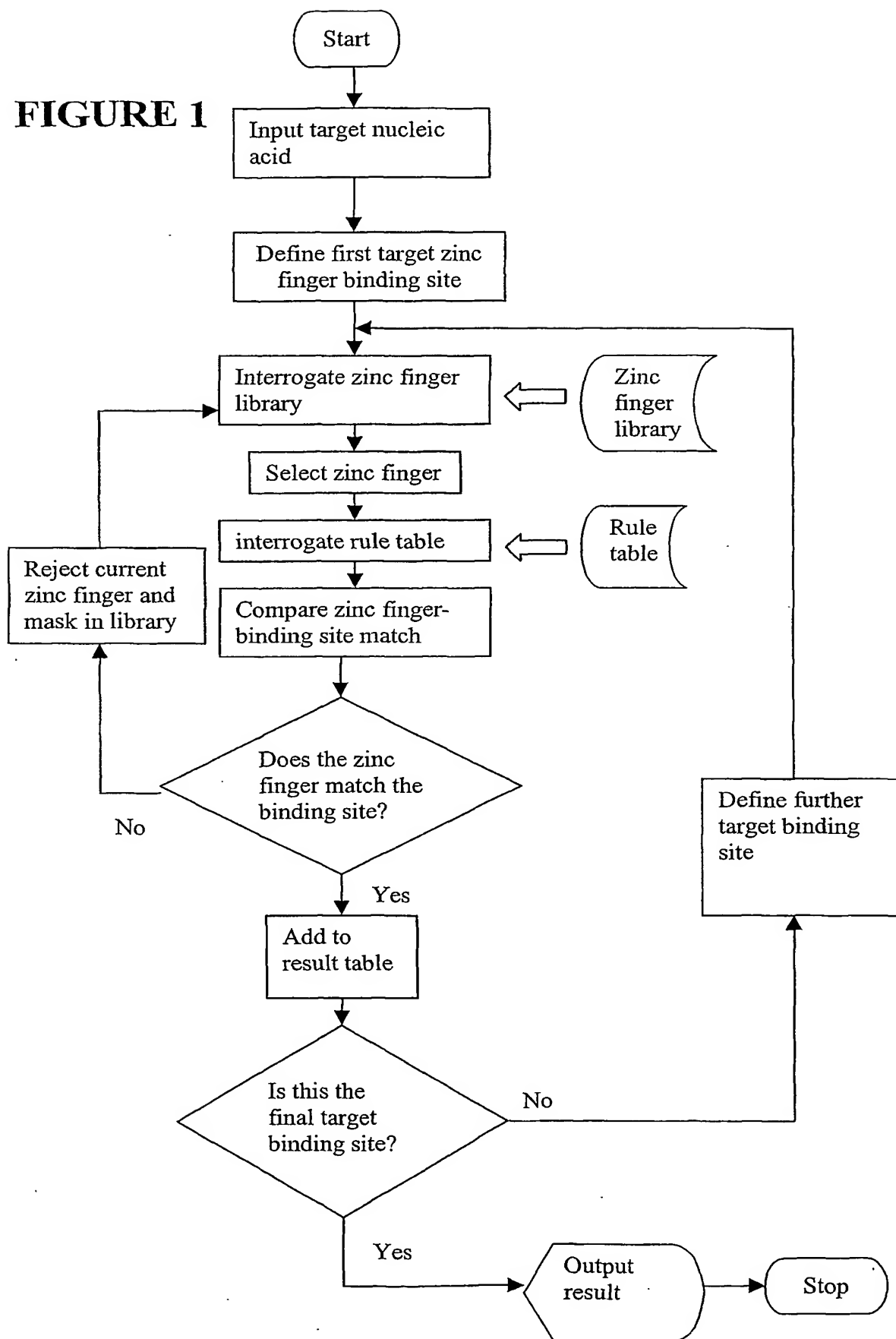
25. A method according to claim 24, further comprising sending instructions to an automated chemical synthesis system to assemble a zinc finger polypeptide as defined by the zinc finger data obtained in (f).

26. A method according to claim 25, wherein the zinc finger polypeptide is tested for binding to the target site, and data from said testing is used to select, from a plurality of candidates, a zinc finger polypeptide capable of binding to the target site.

27. A method according to any one of claims 24 to 26, wherein two or more zinc finger polypeptides are combined to form a zinc finger polypeptide capable of binding to an aggregate binding site comprising two or more target sites.

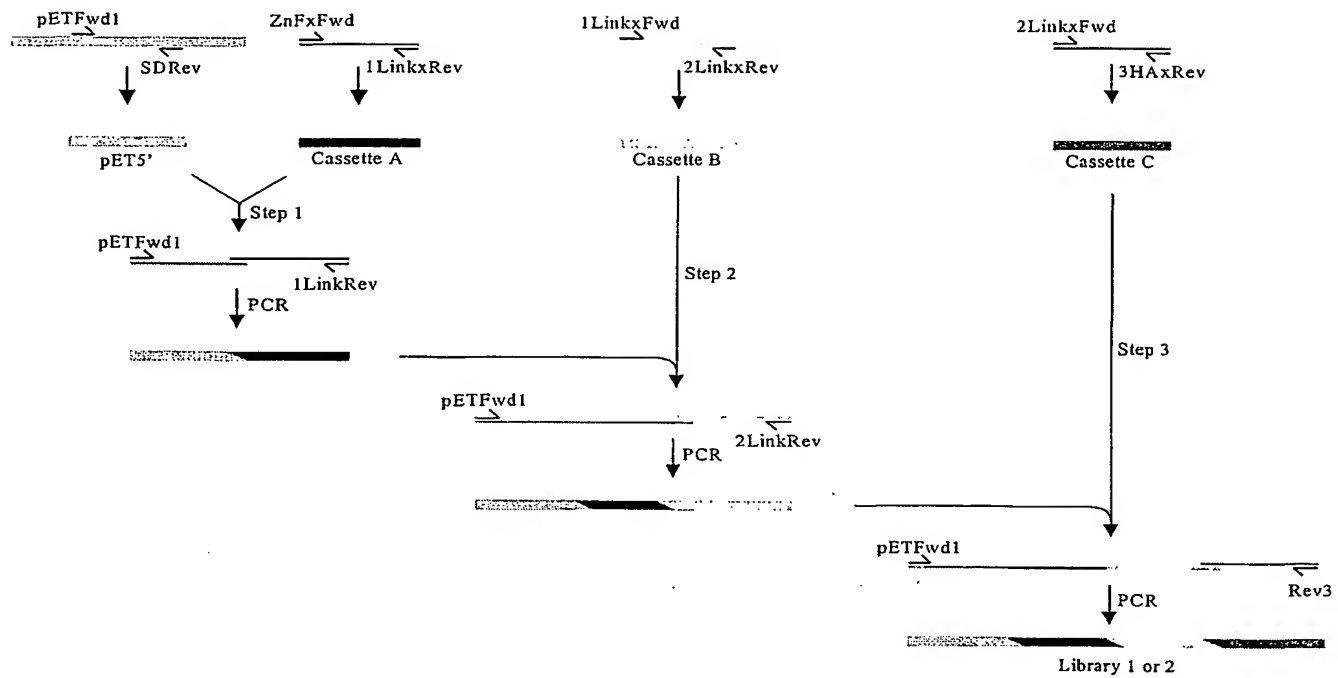
27. A method according to claim 24, wherein the rule table comprises rules as set forth in any one of claims 21 to 23.

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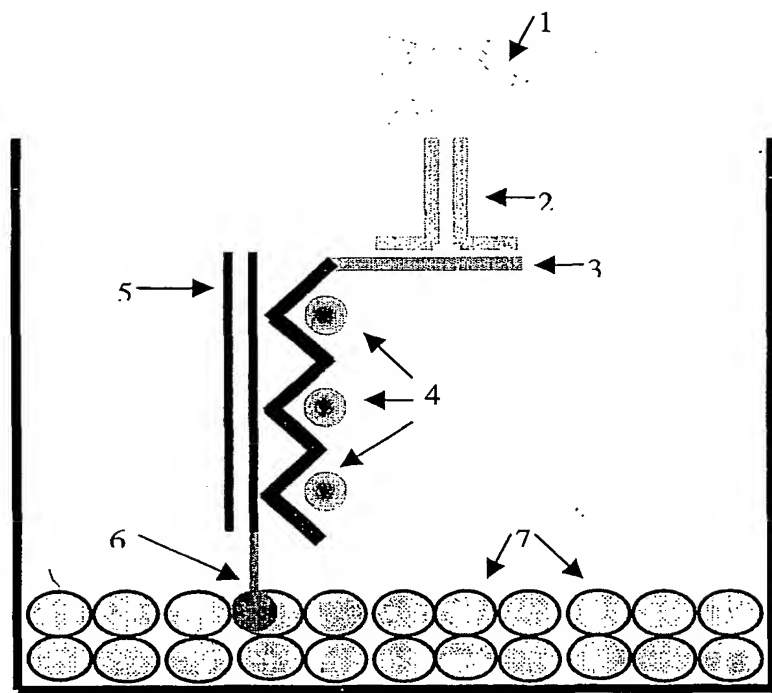
FIGURE 1

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FIGURE 2



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FIGURE 3

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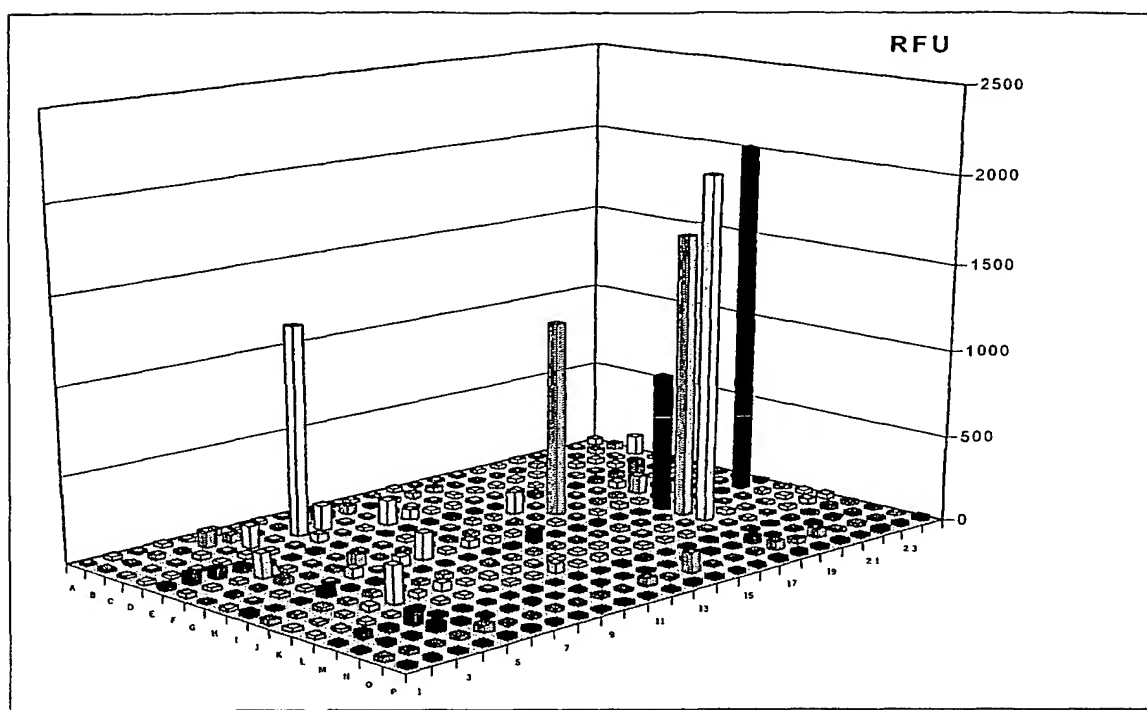
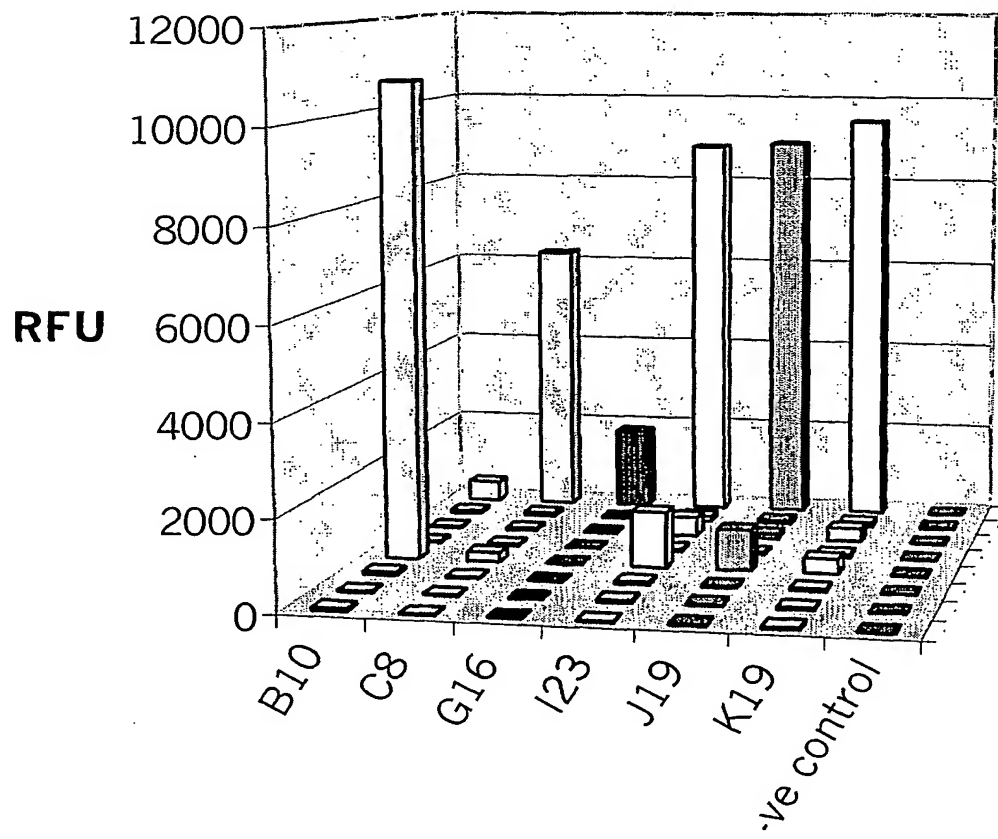


FIGURE 4

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FIGURE 5



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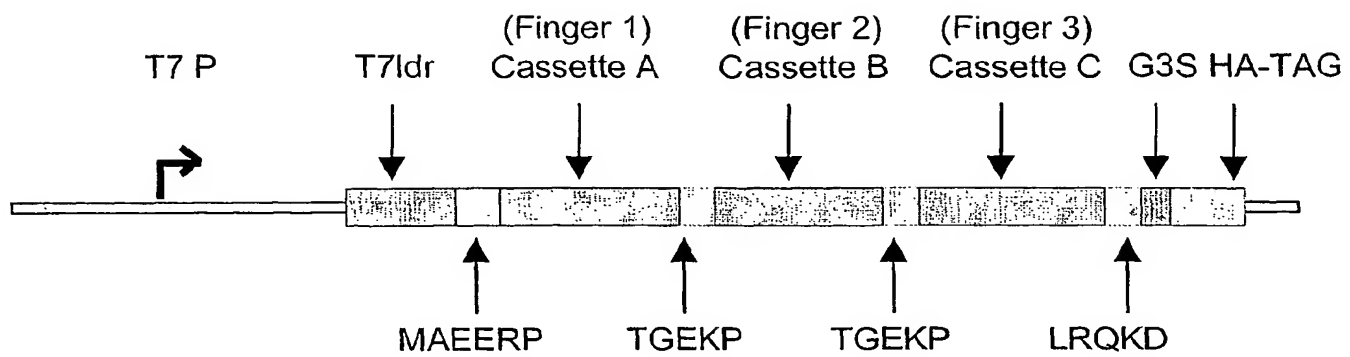


FIGURE 6

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